Final

Treatability Study to Evaluate Remediation by Natural Attenuation at UST Sites 10-494, 10-495, And 10-496
VOLUME 1 OF 1: TEXT AND APPENDICES



BEALE AIR FORCE BASE CALIFORNIA

Prepared for

Air Force Center For Environmental Excellence Technology Transfer Division Brooks Air Force Base San Antonio, Texas

and

9 CES/CEVR Beale Air Force Base, California

March 1999

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FINAL

TREATABILITY STUDY TO EVALUATE REMEDIATION BY NATURAL ATTENUATION AT UST SITES 10-494, 10-495, AND 10-496

at

BEALE AIR FORCE BASE, CALIFORNIA

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LIST OF ABBREVIATIONS

2-D two-dimensional

ADIP alternating-direction implicit procedure

AFB Air Force Base

AFCEE Air Force Center For Environmental Excellence
ASCII American Standard Code for Information Interchange

atm-m³/mol atmosphere-cubic meters per mole

bgs below ground surface
BOD biological oxygen demand

BTEX benzene, toluene, ethylbenzene, and total xylenes

°C degrees centigrade (Celsius)

CaCO₃ calcium carbonate
cfs cubic feet per second
cm/s centimeter per second
DO dissolved oxygen

EAL Evergreen Analytical Laboratory, Inc.

EC electrical conductivity
ET evapotranspiration
°F degrees Fahrenheit

 Fe^{+2} ferrous iron ft foot/feet square feet ft/s feet per second

g/cm³ grams per cubic centimeter HDPE high-density polyethylene

Hg mercury

ID inside diameter

 $\begin{array}{ll} IDW & investigation-derived \ waste \\ K_{oc} & soil \ sorption \ coefficient \end{array}$

kg kilogram L liter

LAW Law Environmental, Inc.

LLNL Lawrence Livermore National Laboratory

LNAPL light nonaqueous-phase liquid

LTM long-term monitoring

MCL Maximum Contaminant Level

μg micrograms mg milligrams

MOC Method of Characteristics

MP monitoring point msl mean sea level mV millivolts

LIST OF ABBREVIATIONS (continued)

MW monitoring well

N nitrogen

NRMRL National Risk Management Research Laboratory

OD outside diameter

ORD Office of Research and Development

OSWER Office of Solid Waste and Emergency Response

Parsons ES Parsons Engineering Science, Inc.

PEI Phillippi Engineering, Inc.

pH negative logarithm of the hydrogen ion concentration

POC point-of-compliance

ppmv parts per million, by volume

PVC polyvinyl chloride QC quality control

r² coefficient of determination
R retardation coefficient
RAOs remedial action objectives
redox reduction/oxidation potential
RI Remedial Investigation

RI/FS Remedial Investigation/Feasibility Study

RMS root mean square

RNA remediation by natural attenuation

RWQCB Regional Water Quality Control Board (California)

second

SAP sampling and analysis plan

SB soil boring
SVP soil vapor point
SW surface water

SWRCB State Water Resources Control Board (California)

TEMB tetramethylbenzene
TMB trimethylbenzene
TOC total organic carbon

TPH total petroleum hydrocarbons

TPH d TPH as diesel fuel TPH-g TPH as gasoline TS treatability study

TVH total volatile hydrocarbons

USEPA United States Environmental Protection Agency

USGS United States Geological Survey

UST underground storage tank VOCs volatile organic compounds

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EXECUTIVE SUMMARY

This report presents the results of a treatability study performed by Parsons Engineering Science, Inc. (Parsons ES) at UST Sites 10-494, 10-495, and 10-496 (UST Site), Beale Air Force Base. California to evaluate the use of remediation by natural attenuation (RNA) with long-term monitoring (LTM) as a remedial option for dissolved benzene, toluene, ethylbenzene, and xylene (BTEX) and total petroleum hydrocarbons as gasoline (TPH-g) contamination in the shallow groundwater. This study focused on the impact of dissolved BTEX and TPH-g compounds on the shallow groundwater system at the site. Site history and the results of soil and groundwater investigations conducted previously are also summarized in this report.

Comparison of BTEX, TPH-g, electron acceptor, and biodegradation byproduct isopleth maps for the UST Site provides strong qualitative evidence of biodegradation of BTEX and TPH-g compounds. Geochemical data strongly suggest that biodegradation of fuel hydrocarbons is occurring at the site via the anaerobic processes of denitrification and iron reduction. Aerobic biodegradation also is likely to occur in the near future due to recent changes in site conditions.

An important component of this study was an assessment of the potential for contamination in groundwater to migrate from the source areas to potential receptors. In particular, this component focused on the contaminant mass loading resulting from the discharge of contaminated groundwater into the surface water drainage channel southeast of the site. To help estimate mass loading rates into the drainage channel and make predictions for future contaminant reductions in groundwater, the Bioplume II model was used to estimate the rate and direction of dissolved BTEX movement through the shallow saturated zone under the influence of advection, dispersion, sorption, and biodegradation. Input parameters for the Bioplume II model were obtained from existing site characterization data, supplemented with data collected by Parsons ES. Extensive site-specific data were used for model implementation. Model parameters that were not measured at the site were estimated using reasonable literature values.

The results of this demonstration suggest that RNA of BTEX and TPH-g compounds is occurring at the UST Site, and that dissolved contamination present in groundwater poses no significant risk to human health or the environment in its present, or predicted future, concentrations and distribution. Model results indicate that eventually the contaminant plume will disappear and that restrictions on groundwater use could be removed in the future. Although discharges of BTEX and TPH-g to surface water have likely occurred in the past, the concentrations of BTEX and TPH-g in groundwater currently discharging to the drainage channel do not appear to be resulting in any detectable contamination in surface water, as confirmed by analysis of surface water samples collected during this study.

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The Air Force recommends that RNA, LTM, and institutional controls be implemented at the site. Bioplume II modeling results suggest that such an approach could result in reduced maximum contaminant concentrations in the source area and reduced concentrations downgradient from the existing plume over time. Institutional controls, such as proper management and disposal of any extracted groundwater and requirements for implementation of adequate health and safety plans for site workers, would protect human health and the environment while remediation is occurring and could be removed in the future once groundwater contaminant concentrations reach water quality objectives.

To verify the Bioplume II model predictions, the Air Force recommends using three LTM wells and three surface water locations to close existing data gaps in the site characterization and monitor the long-term migration and attenuation of the contaminant plume. The presence of free product/mobile LNAPL should be evaluated during installation of the new LTM well in the source area and removed to the extent practicable in compliance with regulatory requirements. One of the new LTM wells should be installed on the eastern side of the drainage channel to confirm that the groundwater contaminants have not migrated beneath the drainage channel. Because of the removal of the compost pile near the source area at this site, the groundwater geochemistry can be expected to change within the next few years. Therefore, it is recommended that the assimilative capacity of site groundwater be reevaluated during LTM.

Groundwater and surface water samples and elevation measurements should be collected quarterly for at least one year to determine seasonal variations in groundwater flow direction and gradient and contaminant concentrations at the site. Thereafter, it is expected that sampling frequency could be reduced to annually. The LTM plan presented in this report will serve as an interim LTM until installation of the LTM wells and completion of the first year of sampling (4 quarterly sampling rounds). Regular sampling and analysis of groundwater and surface water from LTM wells and surface water locations will allow the effectiveness of RNA to be monitored and should allow sufficient time to implement additional engineering controls to contain the plume if BTEX or TPH-g compounds are detected above levels of concern in the surface water or increase significantly in the groundwater.

Contaminant fate and transport model results indicate that LTM wells and surface water locations should be sampled for 15 years, after which time it is expected that the plume will disappear. Both groundwater samples and surface water samples should be analyzed for BTEX compounds by US Environmental Protection Agency (USEPA) Method SW8021 and TPH-g by USEPA Method 8015B. Groundwater samples should also be analyzed for dissolved oxygen and additional geochemical parameters to verify the progress of RNA. If any BTEX or TPH-g concentrations detected in surface water exceed the water quality objectives or concentrations in groundwater are increasing significantly greater than that predicted by modeling results, additional corrective actions and comparative cost analyses may be required to remediate groundwater at the site.

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SECTION 1

INTRODUCTION

This report was prepared by Parsons Engineering Science, Inc. (Parsons ES) and presents the results of a treatability study (TS) conducted to evaluate the use of remediation by natural attenuation (RNA) for remediation of groundwater contaminated by petroleum hydrocarbons at UST Sites 10-494, 10-495, and 10-496 (UST Site), Beale Air Force Base (AFB), California. Previous investigations have shown that soils and groundwater at this site are contaminated with gasoline.

As used throughout this report, RNA refers to a management strategy that relies on natural attenuation mechanisms to remediate contaminants dissolved in groundwater, to control receptor exposure risks associated with contaminants in the subsurface, and to meet water quality objectives. The United States Environmental Protection Agency (USEPA) Office of Research and Development (ORD) and Office of Solid Waste and Emergency Response (OSWER) define natural attenuation as:

The biodegradation, dispersion, sorption, volatilization, and/or chemical and biochemical stabilization of contaminants to effectively reduce contaminant toxicity, mobility, or volume to levels that are protective of human health and the ecosystem.

As suggested by this definition, mechanisms for natural attenuation of BTEX include advection, dispersion, dilution from recharge, sorption, volatilization, and biodegradation. Of these processes, biodegradation is the only mechanism working to transform contaminants into innocuous byproducts. Contaminant destruction occurs through biodegradation when indigenous microorganisms work to bring about a reduction in the total mass of contamination in the subsurface without the addition of nutrients. Patterns and rates of natural attenuation can vary markedly from site to site depending on governing physical and chemical processes.

The main emphasis of the work described herein was to evaluate the potential for naturally occurring degradation mechanisms to reduce dissolved BTEX concentrations in groundwater to levels that are protective of human health and the environment.

1.1 SCOPE AND OBJECTIVES

Parsons ES, in conjunction with researchers from the United States Environmental Protection Agency (USEPA) National Risk Management Research Laboratory (NRMRL) Subsurface Protection and Remediation Division, was retained by the United States Air Force Center for Environmental Excellence (AFCEE) Technology Transfer Division to conduct site

characterization and groundwater modeling to evaluate the scientific defensibility of RNA with long-term monitoring (LTM) as a remedial option for gasoline-contaminated groundwater at the UST Site at Beale AFB. Performance of numerous tasks was required in order to fulfill the project objective. These tasks included:

- Reviewing existing hydrogeologic and soil and groundwater quality data for the site;
- Conducting supplemental site characterization activities to determine the nature and extent of soil, groundwater, and surface water contamination;
- Collecting geochemical data in support of RNA;
- Developing a conceptual hydrogeologic model of the shallow saturated zone, including the current distribution of contaminants;
- Evaluating site-specific data to determine whether naturally occurring processes of contaminant attenuation and destruction are occurring in groundwater at the site;
- Designing and executing a Bioplume II model for site hydrogeologic conditions:
- Simulating the fate and transport of fuel hydrocarbons in groundwater under the influence of advection, dispersion, adsorption, and biodegradation using the Bioplume II model;
- Evaluating a range of model input parameters to determine the sensitivity of the model to those parameters and to consider, if necessary, several contaminant fate and transport scenarios;
- Determining if naturally occurring processes are sufficient to minimize BTEX plume expansion so that groundwater and surface water quality objectives can be met at a downgradient point of compliance (POC);
- Conducting a preliminary exposure pathways analysis for potential current and future receptors:
- Developing remedial action objectives (RAOs) and reviewing available remedial technologies;
- Using the results of modeling to recommend the most appropriate remedial option based on specific effectiveness, implementability, and cost criteria; and
- Providing an LTM plan that includes LTM and POC locations and a sampling and analysis plan (SAP).

Site characterization activities in support of RNA conducted for this report included soil sampling for lithological characterization; installation of groundwater monitoring points; static groundwater level measurement at existing monitoring wells and newly installed monitoring points; groundwater sample collection and analysis from existing site monitoring wells and newly installed monitoring points; and surface water sample collection and

analysis from two drainage ditches downgradient from the site. Previous site investigation activities and results were used to guide placement of sampling locations for this study.

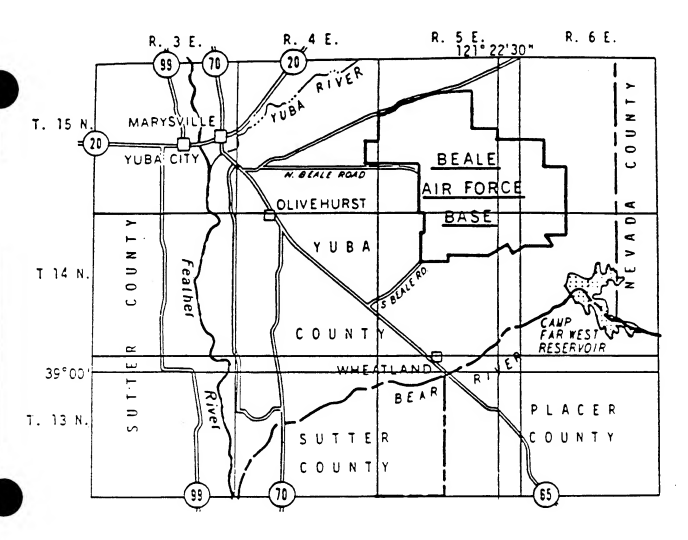
Site-specific data were used to develop groundwater flow and solute fate and transport models for the site using Bioplume II, and to conduct a preliminary receptor exposure pathways analysis. The modeling effort was used to predict the future extent and concentrations of the dissolved contaminant plume by modeling the combined effects of advection, dispersion, sorption, and biodegradation on dissolved BTEX. Results of the model were used to assess the potential for completion of exposure pathways involving groundwater and to determine whether RNA with LTM and institutional controls is an appropriate and defensible remedial option for contaminated groundwater. The results will be used to provide technical support for the RNA with LTM and institutional controls remedial option during regulatory negotiations, as appropriate.

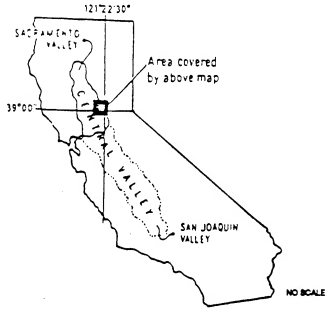
Alternate remedial options were considered to identify the major advantages and disadvantages associated with different groundwater remedial strategies. These options included groundwater extraction/containment via vertical wells and/or interceptor trenches (pump and treat); air sparging; physical containment using slurry walls, grout curtains, or sheet piles; and funnel and reactive gate walls. Hydrogeologic and groundwater chemical data necessary to evaluate these remedial options were either collected under this program or were available from previous investigations of this site or from the technical literature. Field work conducted under this program, however, was oriented toward the collection of hydrogeologic data to be used as input into the Bioplume II groundwater model in support of RNA with LTM for restoration of fuel-hydrocarbon-contaminated groundwater.

This TS contains nine sections, including this introduction, and six appendices. Section 2 summarizes site characterization activities. Section 3 summarizes the physical characteristics of the study area. Section 4 describes the nature and extent of soil, groundwater, and surface water contamination and the geochemistry of groundwater at the site. Section 5 describes the Bioplume II model and design of the conceptual model for the site; lists model assumptions and input parameters; and describes sensitivity analysis, model output, and the results of the Bioplume II modeling. Section 6 presents a comparative analysis of remedial alternatives. Section 7 presents the LTM plan for the site. Section 8 presents the conclusions of this work and provides recommendations for further work at the site. Section 9 lists the references used to develop this document. Appendix A contains borehole logs and monitoring point construction diagrams. Appendix B contains analytical results from previous site investigations. Appendix C presents soil, groundwater, and surface water analytical results from this study. Appendix D contains model input parameters and calculations related to model calibration. Appendix E contains Bioplume II model input and output in American Standard Code for Information Interchange (ASCII) format on a diskette. Appendix F contains calculations for remedial option design and costing.

1.2 FACILITY BACKGROUND

Beale AFB is located in Yuba County, California, approximately 50 miles north of Sacramento and 10 miles east of the city of Marysville (Figure 1.1). The Base occupies approximately 23,000 acres of land in Yuba County and has 8,000 military and non-military





SOURCE: GROUNDWATER CONDITIONS AT BEALE AIR FORCE BASE AND VICINITY, CALIFORNIA U.S. GEOLOGICAL SURVEY OPEN-FILE REPORT 80-204, PAGE 1980



0 2 4

APPROX. SCALE IN MILES

FIGURE 1.1

VICINITY MAP

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California



PARSONS ENGINEERING SCIENCE, INC.

Alameda, California

\722450\14\GRAPHICS\FIGS P8.CDR 10/28/96

personnel on site on a daily basis. The land surrounding the Base is primarily used for agriculture.

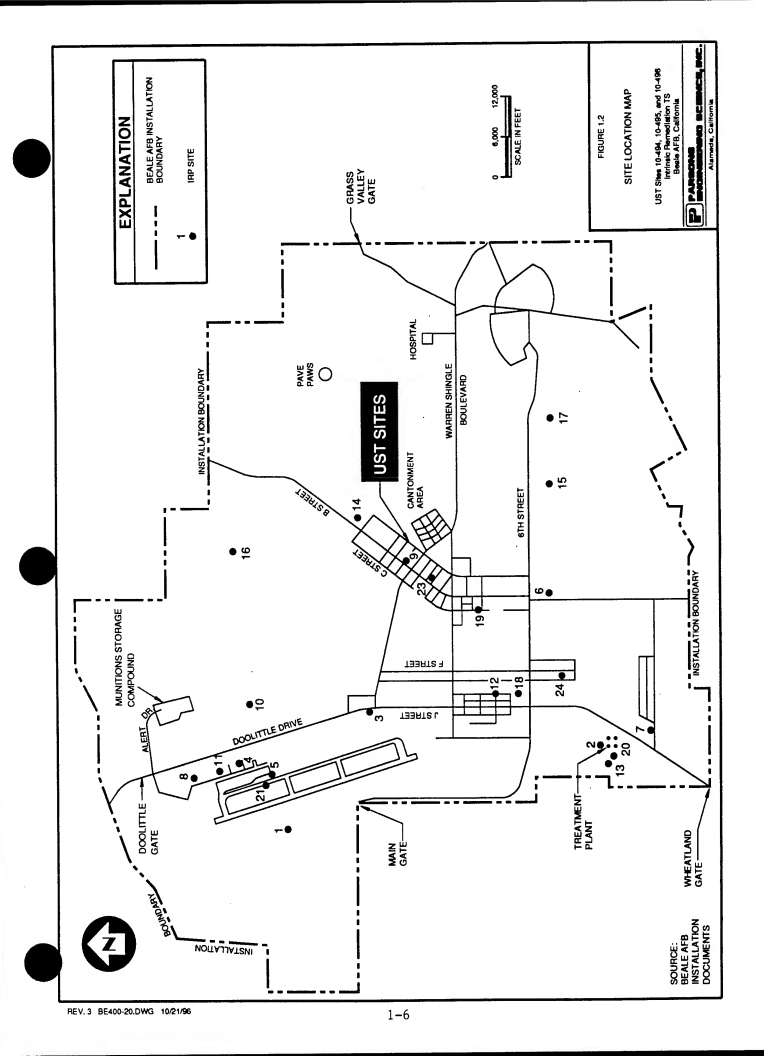
The project site is located in the central portion of the Base (Figure 1.2) near the intersection of 30th and A Streets (Figure 1.3). Three 12,000-gallon USTs (which have been designated USTs 10-494, 10-495, and 10-496) were installed at the site in the early 1940s. The USTs were placed on concrete saddles set 6 feet below ground surface (bgs). The tanks were used primarily for gasoline storage and were part of a series of six armored tank division fueling stations located along A Street. All three tanks are reported to have been removed prior to 1962; however, the preformed concrete saddles were left in place. The fuel pipelines that supplied the USTs are inactive but still in place. Law Environmental, Inc. (LAW) and Metcalf & Eddy conducted investigations of the pipeline, which runs throughout the Base, in 1994 (LAW, 1995).

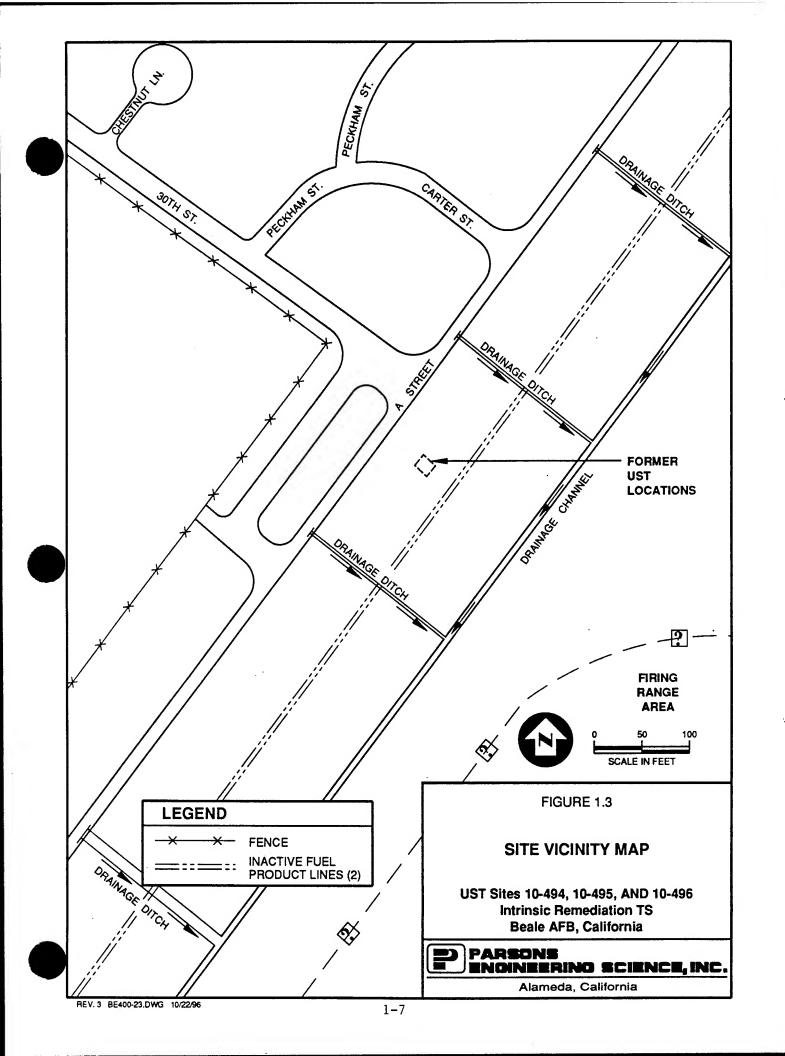
The excavation pit, which was left open after tank removal activities, was used as a compost pile for landscape waste until November 1995. In November 1995, the compost pile was removed, the excavation pit was filled with clean soil, and the site was completely regraded. There are no buildings in the immediate vicinity of the site, and the land surrounding the site is not currently in use. The site is surrounded on three sides by surface water drainage ditches; however, only the largest drainage channel east of the site contains water year-round.

In December 1994, LAW (1995) conducted a preliminary site investigation that included advancement of seven soil vapor points to depths ranging from 5.5 to 8.5 feet bgs. In February 1995, Metcalf & Eddy advanced seven soil borings around the site to depths ranging from 6 to 18 feet bgs. Two of these soil boreholes, SB-1 and SB-2, were converted to monitoring wells, designated MW-1 and MW-2. In addition, a total of seven test pits were excavated to depths between 6 and 9 feet bgs.

Groundwater was encountered in soil boreholes and test pits at depths of 3 to 6 feet bgs. During the drilling of boreholes and vapor points in the immediate vicinity of the tank pit area. a visual inspection of groundwater indicated the presence of mobile light nonaqueous-phase liquid (LNAPL), or free product. The full extent of contamination, the source of contamination, and the groundwater flow direction and gradient were not determined during prior investigations. Analytical results from soil, groundwater, and soil vapor sampling conducted during the prior investigations are discussed in Section 4 and detailed in Appendix B.

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SECTION 2

SITE CHARACTERIZATION ACTIVITIES

This section presents the methods used by Parsons ES to collect additional site-specific data at Beale AFB. To meet the requirements of the RNA demonstration, additional data were required to evaluate the near-surface geology and hydrogeology, aquifer properties, and the nature and extent of contamination at the UST Site. Site activities included installing temporary groundwater monitoring points (MPs); collecting and analyzing soil, groundwater, and surface water samples; and measuring and estimating hydrogeologic parameters (static groundwater levels, groundwater gradient, groundwater flow direction, and hydraulic conductivity). The physical and chemical hydrogeologic parameters determined during field work activities included:

- Depth from measurement datum to the water table or potentiometric surface in existing monitoring wells and new monitoring points;
- Stratigraphic analysis of subsurface media:
- Rate of change of water elevation following rapid depression or elevation of water level in a monitoring well;
- Locations of potential groundwater recharge and discharge areas:
- Extent of residual petroleum hydrocarbon contamination in soils;
- Additional chemical analysis of soil, groundwater, and surface water for the parameters listed in Table 2.1.

Previously collected data and data collected under this program were integrated to aid with interpretation of the physical setting (Section 3) and contaminant distribution (Section 4), and to develop the conceptual site model (Section 5).

Twenty-two (22) temporary groundwater MPs were installed to further characterize the site and determine the extent of groundwater contamination. Procedures used to install the MPs and to collect soil core samples are described in Section 2.1. Procedures used to sample existing groundwater monitoring wells and newly installed temporary groundwater MPs are described in Section 2.2. Procedures used to sample surface water are described in Section 2.3. Sample handling procedures are described in Section 2.4.

Aquifer testing was performed during this investigation. Procedures used to conduct the aquifer slug tests are presented in Section 2.5. The calculated hydraulic conductivities from

TABLE 2.1 ANALYTICAL PROTOCOLS FOR GROUNDWATER, SURFACE WATER, AND SOIL SAMPLES

UST Sites 10-494, 10-495, and 10-496

Intrinsic Remediation TS Beale AFB, California

MATRIX	METHOD	FIELD (F) OR ANALYTICAL LABORATORY (L)
GROUNDWATER AND SURFACE	WATER *	
Total Iron *	Colorimetric, HACH Method 8008	F
Ferrous Iron (Fe ²⁺) *	Colorimetric, HACH Method 8146	F
Ferric Iron (Fe ³⁺) *	Difference between total and ferrous iron	F
Manganese *	Colorimetric, HACH Method 8034	F
Sulfide *	Colorimetric, HACH Method 8131	F
Carbon Dioxide *	CHEMetrics Method 4500	F
Alkalinity (Carbonate [CO3-2] and Bicarbonate [HCO3-1]) *	Titrimetric, HACH Method 8221	F
Redox Potential	A2580B, direct reading meter	F
Dissolved Oxygen	E360.1, direct reading meter	F
рН	E150.1/SW9040, direct reading meter	F
Conductivity	E120.1/SW9050, direct reading meter	F
Temperature	E170.1, direct reading meter	F
Nitrate	E300.0	L
Nitrite	E300.0	L
Chloride	E300.0	L
Sulfate	E300.0	L
Alkalinity	E310.1	L
Methane *	RSKSOP-175	L
Total Organic Carbon *	E415.1	L
Aromatic Hydrocarbons	SW8020A	1
(including Trimethylbenzene)		L
Total Petroleum Hydrocarbons, as		
gasoline and diesel	SW8015M	L
AmmoniaDiss. Gas in Water *	E350.3	L
SOIL		
Total Organic Carbon	SW9060	L
Grain-size distribution	ASTM-D422	L
Moisture	ASTM D-2216	L
Aromatic Hydrocarbons	SW8020A	L
Total Petroleum Hydrocarbons, as		
gasoline and diesel	SW8015M	L

^{*} Performed for groundwater samples only

the aquifer tests were used to estimate the velocity of groundwater and contaminants in the shallow saturated zone, and were used as input to the Bioplume II model. The calculated range of hydraulic conductivity values was used to guide the model sensitivity analysis. The remainder of Section 2 describes the procedures followed during TS field work activities. Additional details regarding investigative techniques are presented in the work plan for the site (Parsons ES, 1995).

2.1 MONITORING POINT INSTALLATION AND SOIL SAMPLE COLLECTION

Twenty-two (22) temporary groundwater MPs were installed using the Geoprobe[®] system in February/March 1996. The truck-mounted Geoprobe[®] system is a probe-driven sampler and is described in detail in Section 2.1.1.1. Soil samples were collected for lithologic description and laboratory analysis during installation of the MPs. Existing monitoring wells and the temporary MPs are shown on Figure 2.1.

Although the work plan called for four proposed MPs on the southeastern side of the main northeast/southwest drainage channel, site access problems at the time of field work prevented installation of these MPs. Two additional MPs called for in the work plan were not installed because field conditions indicated that the groundwater upgradient from the site could be adequately characterized with fewer MPs.

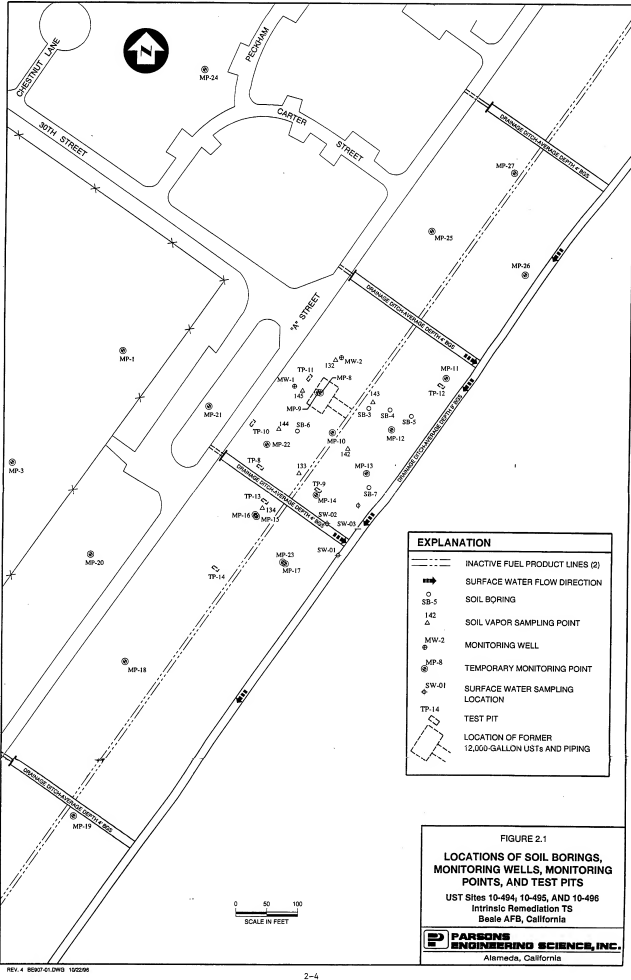
Nested MP clusters (i.e., multiple points installed at the same location and screened at different depths) were installed at three locations (MP-8/MP-9, MP-15/MP-16, and MP-17/MP-23). All shallow MPs were screened at the top of the shallow aquifer between approximately 5 and 10 feet bgs, generally in silty and/or gravelly fine- to coarse-grained sands. The deeper MPs at the nested MP clusters were constructed of stainless-steel screens. approximately 6 inches in length, and installed at approximately 17 feet bgs in the underlying clayey silt.

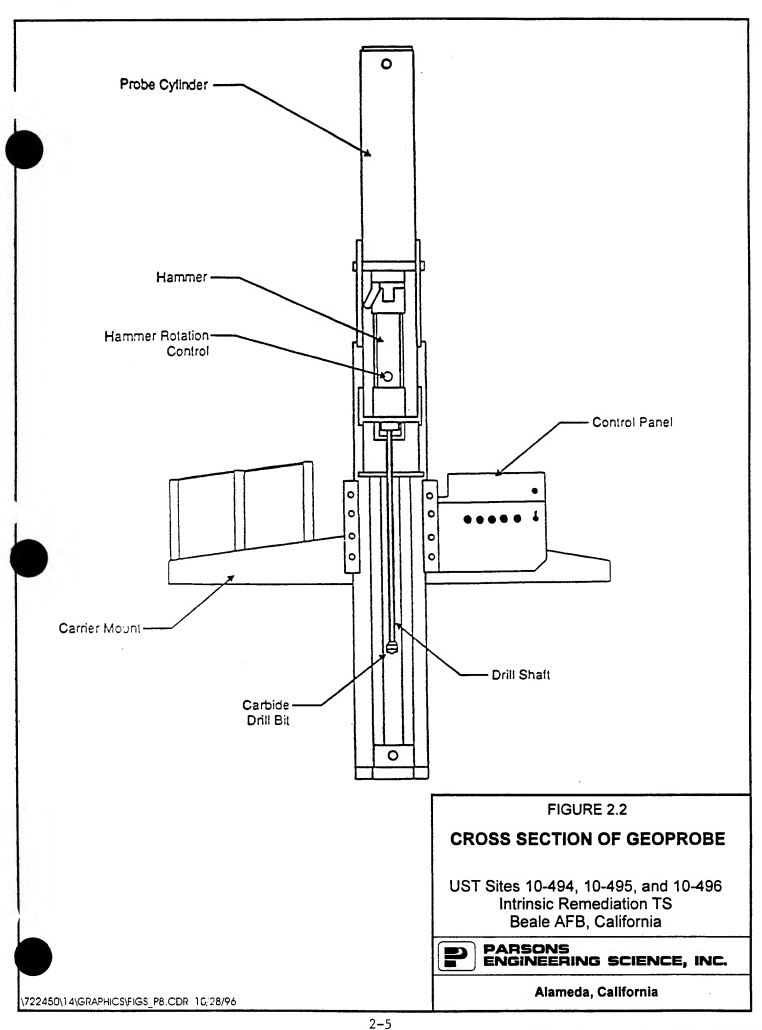
2.1.1 Installation Procedures and Determination of Stratigraphy

2.1.1.1 Geoprobe® Direct-Push Technology

The Geoprobe[®] system is a hydraulically powered percussion/probing machine used to advance sampling tools through unconsolidated soils. This system provides for the rapid collection of soil, soil gas, and groundwater samples at shallow depths while minimizing the generation of investigation-derived waste (IDW) materials. Figure 2.2 is a diagram of the Geoprobe[®] system.

The probe-drive sampler serves as both the driving point and the sample collection device. and is attached to the leading end of the probe rods. To collect a soil sample, the sampler is pushed or driven to the desired sampling depth, the drive point is retracted, which opens the sampling barrel, and the sampler is subsequently pushed into the undisturbed soils. The soil cores are retained within brass, stainless steel, or clear acetate liners inside the sampling barrel. The probe rods are then retracted, bringing the sampling device to the surface. Soil





samples can then be extruded from the liners for logging or, when required, the liners can be capped, and undisturbed samples can be submitted to the analytical laboratory for testing.

2.1.1.2 Determination of Stratigraphy

During installation of the MPs, a Parsons ES field scientist observed all field investigation activities and maintained a detailed descriptive log of all subsurface materials recovered during soil coring. The boring logs for the 22 MPs are included in Appendix A, and results are discussed in Section 3 and Section 4.

2.1.1.3 Geoprobe® Equipment Decontamination

Prior to arriving at the site, probe rods, tips, sleeves, pushrods, samplers, tools, and other downhole equipment was decontaminated using a high-pressure, steam/hot water wash. Between sampling locations, equipment was decontaminated with a phosphate-free detergent and a potable water rinse, and allowed to air dry. Only potable water was used for decontamination.

All rinse water used for decontamination was placed in a temporary holding tank located northwest of the site. A low volume of soil cuttings (sampling waste) was generated during MP installation. Disposal of the soil cuttings and wastewater is discussed in Section 2.2.1.1.

2.1.2 Monitoring Point Installation and Construction

The 19 shallow MPs were installed by punching and sampling a borehole with the Geoprobe[€]. Upon removing the rods, the borehole depth was measured to determine if the hole remained open. All shallow MPs installed at the UST Site remained open after withdrawing the rods; therefore, the PVC casing and screen were placed at the appropriate depths, the annular space around the screen was filled with sand filter pack, and the annulus around the casing was filled with bentonite. The 19 shallow MPs were constructed using a 0.5-inch inside-diameter (ID)/0.75-inch outside-diameter (OD) polyvinyl chloride (PVC) screen and casing. Installed screens were 5 feet in length and factory-slotted with 0.010-inch openings. A sacrificial tip, screwed into the PVC screen, served as the bottom cap for the MP.

The 3 deep MPs in the nested MP clusters (MP-8, MP-16, and MP-23) were installed by punching and sampling a borehole to several feet above the target depth. The probe rods were withdrawn from the borehole, and the soil sampler was replaced with the well point assembly. An appropriate length of Teflon® tubing was threaded through the probe rods and attached to the well point. The assembly was lowered into the borehole and then driven down to the target depth. The probe rods were removed, leaving the sacrificial tip. screen assembly, and tubing behind. The annular space around the tubing was filled with granular bentonite. The 3 deep MPs were constructed using a sacrificial drive point attached to a 6-inch length of 0.5-inch-diameter stainless steel mesh screen, which was in turn connected to 0.375-inch Teflon® tubing.

Well materials were inspected for cleanliness prior to use. No glue or solvents were used. Construction details were noted on a Monitoring Point Installation Record form and are summarized in Appendix A.

2.1.3 Development

The MPs were developed prior to sampling to remove fine sediments from the portion of the formation adjacent to the MP screens. Development was accomplished using a peristaltic pump with new, dedicated high-density polyethylene (HDPE) tubing. The pump tubing was regularly lowered to the bottom of the MP so that fines were agitated and removed from the MP in the development water.

When a constant flow of groundwater could be obtained from an MP, development was continued until the groundwater was relatively free of fine sediments. Water was removed until pH, temperature, and water clarity (turbidity) stabilized. Monitoring point development occurred a minimum of 24 hours prior to sampling. All development waters and purge waters were handled as detailed in Section 2.2.1.1.

2.1.4 Site Restoration and Monitoring Point Abandonment

After the MPs were installed and sampled, each location was restored as closely to its original condition as possible. All MPs were abandoned by sealing with a bentonite-cement grout hydrated with potable water to eliminate any creation or enhancement of contamination migration pathways to the groundwater.

2.2 GROUNDWATER SAMPLING

Groundwater samples were collected at the 2 existing monitoring wells (MW-1 and MW-2) and 18 of the 22 newly-installed MPs (Figure 2.1). A total of 4 MPs — MP-1, plus all 3 of the deep MPs (MP-8, MP-16, and MP-23) — could not be sampled because a sufficient volume of groundwater could not be obtained.

Groundwater samples were analyzed by Parsons ES field scientists for dissolved oxygen (DO), reduction/oxidation potential (redox), total iron, ferrous iron, soluble manganese, free carbon dioxide, temperature, pH and electrical conductivity. Analyses for total petroleum hydrocarbons (TPH), and volatile aromatics (including BTEX) were performed by Curtis and Tompkins of Berkeley, California. Analyses for nitrate, nitrite, sulfate, methane, alkalinity, dissolved ammonia, chloride, and total organic carbon (TOC) were performed by Evergreen Analytical Laboratory, Inc. (EAL) of Wheatridge, Colorado.

Groundwater sampling forms were used to document the specific details of the sampling event for each well and MP. Groundwater sampling results from the analytical laboratory are included as Appendix C. Results for groundwater sampling previously conducted are included in Appendix B.

This section describes the procedures used for collecting groundwater samples. In order to maintain a high degree of quality control (QC) during this sampling event, the procedures

described in the site work plan (Parsons ES, 1995) and summarized in the following sections were followed.

2.2.1 Preparation for Sampling

All equipment to be used for sampling was assembled and properly cleaned and calibrated (if required) prior to arriving in the field. Special care was taken to prevent contamination of the groundwater and extracted samples. The two primary ways in which sample contamination can occur are through contact with improperly cleaned equipment and through cross-contamination due to insufficient cleaning of equipment between wells and monitoring points. To prevent such contamination, downhole sampling equipment and the water level probe and cable used to determine static water levels were thoroughly cleaned before and after field use and between uses at different sampling locations according to the procedures presented in Section 2.2.1.1.

2.2.1.1 Equipment Decontamination and Waste Disposal

All portions of sampling and test equipment that were used for sampling or were used downhole were thoroughly cleaned before each use. This included the Geoprobe[®] soil sampling tool, pump tubing, water level probe and cable, lifting line, and other equipment or portions thereof that came into contact with the samples. Based on the types of sample analyses conducted, the following cleaning protocol was used:

- Washed with potable water and phosphate-free laboratory detergent;
- Rinsed with potable water;
- Rinsed with isopropyl alcohol; and
- Rinsed with distilled or deionized water.

When precleaned, disposable sampling equipment was used, the cleaning protocol specified above was not required. Laboratory-supplied sample containers were cleaned and sealed by the laboratory. The type of container provided and the method of container decontamination was documented in the permanent record of the sampling event. Any deviations from these procedures were documented in the field scientist's field notebook and on the groundwater sampling record.

All purge water, decontamination rinseate, and other wastewaters were placed in a temporary holding tank located northwest of the site. Approximately 150 gallons of water was generated during installation of the MPs and sampling activities. After sampling activities were completed, the water was sampled, analyzed, and discharged at the Base treatment pad for contaminated soils located near the intersection of 9th Street and J Street and operated by Metcalf & Eddy. All soil cuttings generated during MP installation also were removed to the Base treatment pad.

2.2.1.2 Equipment Calibration

As required, field analytical equipment was calibrated according to the manufacturers' specifications prior to field use. Calibration records for field analytical equipment were maintained by Parsons ES, and field calibrations were recorded in the field notebook. Calibration was required for equipment used for onsite measurements of DO, pH, electrical conductivity, temperature, redox potential, ferrous iron (Fe²⁺), and other field parameters listed in Table 2.1.

2.2.1.3 Preparation of Location

Prior to proceeding with the sampling, the areas around the existing monitoring wells (MWs) and new MPs were cleared of foreign materials, such as brush, rocks, and debris. These procedures prevented sampling equipment from inadvertently contacting debris around the sampling location. Any irregularities with the protective cover, cap, lock, external surface seal, internal surface seal, well identification, well datum, and pad for existing MWs were noted.

2.2.1.4 Water Level and Total Depth Measurements

Prior to removing any water from the MWs or MPs, the static water level was measured. An electric water level probe was used to measure the depth to groundwater below the datum to the nearest 0.01 foot. After measuring the static water level, the water level probe was slowly lowered to the bottom of the monitoring well/point and the depth was measured to the nearest 0.01 foot. Based on these measurements, the volume of water to be purged from the MW or MP was calculated. Water levels used to develop the site groundwater potentiometric surface map presented in Section 3 were taken at the conclusion of all field activities on 20 March 1996.

2.2.2 Purging and Sample Collection

MW and MP purging consisted of the removal of at least three casing volumes of water prior to sample collection. A peristaltic pump was used for MW and MP purging. Once three casing volumes of water were removed from the MW or MP, purging continued until the temperature, redox potential, electrical conductivity, pH, and DO concentrations had stabilized, and if possible, until the purge water became clear.

When sufficient water was available, sample collection commenced immediately after completion of the purge. In all instances, groundwater samples were collected within 24 hours of the purge. Groundwater samples were extracted with a peristaltic pump. Sample containers were filled directly from the pump discharge tube. The groundwater was directed toward the bottle wall near the top and allowed to run down the inner walls of the sample bottle in order to minimize aeration of the sample. Sample containers were completely filled so that no air space remained in the container.

2.2.3 Onsite Chemical Parameter Measurement

As indicated in Table 2.1, many of the groundwater chemical parameters were measured on site by Parsons ES field scientists. Some of the measurements were made with direct-reading meters, while others were made using a Hach[®] portable colorimeter in accordance with specific Hach[®] analytical procedures. These measurements are briefly summarized in the following sections; more detailed descriptions are provided in the work plan for the TS (Parsons ES, 1995).

2.2.3.1 Dissolved Oxygen Measurements

DO measurements were made during purging and immediately prior to groundwater sample acquisition in a flow-through cell using a direct-reading meter (YSI® model 33). DO concentrations were recorded after each volume was purged. The stable value recorded at the end of the purge was assumed to represent the aquifer characteristics at each location.

2.2.3.2 pH, Temperature, and Specific Conductance

Because the pH, temperature, and specific conductance of a groundwater sample can change significantly within a short time following sample acquisition, these parameters were measured in the field in unfiltered, unpreserved "fresh" water collected by the same technique as the samples taken for laboratory analyses. The measurements were made with direct-reading meters in a flow-through cell or a clean glass container separate from those intended for laboratory analysis, and the measured values were recorded in the groundwater sampling record.

2.2.3.3 Carbon Dioxide Measurements

The presence of free carbon dioxide dissolved in groundwater is unlikely because of carbonate buffering; however, carbon dioxide concentrations in the dissolved contaminant plume should be compared to background concentrations to determine whether they are elevated. Carbon dioxide concentrations in groundwater were measured in the field by Parsons ES scientists via titrimetric analysis using Hach® Method 8223.

2.2.3.4 Total Iron, Ferrous Iron, and Ferric Iron Measurements

Iron is an important trace nutrient for bacterial growth, and different valence states of iron can affect the redox potential of the groundwater and act as an electron acceptor for biological metabolism under anaerobic conditions. Iron concentrations were measured in the field via colorimetric analysis with a Hach® DR/700 Portable Colorimeter after appropriate sample preparation. Hach® Method 8008 for total soluble iron (0 to 3.0 milligrams per liter [mg/L] Fe³⁺ + Fe²⁺) and Hach® Method 8146 for ferrous iron (0 to 3.0 mg/L Fe²⁺) were used to prepare and quantitate the samples. Ferric iron (Fe³⁺) was quantitated by subtracting ferrous iron concentrations from the total iron concentration.

2.2.3.5 Manganese Measurements

Manganese is a potential electron acceptor under anaerobic environments. Manganese concentrations were measured in the field using colorimetric analysis with a Hach[®] DR/700 Portable Colorimeter. USEPA-approved Hach[®] Method 8034 (0 to 20.0 mg/L Mn) was used for quantitation of manganese concentrations.

2.2.3.6 Redox Potential

The redox potential of groundwater is an indication of the relative tendency of a solution to accept or transfer electrons. Redox reactions in groundwater are usually biologically mediated; therefore, the redox potential of a groundwater system depends upon and influences rates of biodegradation. Redox potential can be used to provide real-time data on the location of the contaminant plume, especially in areas undergoing anaerobic biodegradation. The redox potential of a groundwater sample taken inside the contaminant plume should be somewhat less than that taken upgradient or crossgradient from the plume.

The redox potential of a groundwater sample can change significantly within a short time following sample acquisition and exposure to atmospheric oxygen. Therefore, this parameter was measured in the field in a flow-through cell by the same technique used for DO measurements.

2.3 SURFACE WATER SAMPLING

2.3.1 Preparation of Location

Three surface water samples (SW-01 through SW-03, Figure 2.1) were collected from the drainage ditches located downgradient from the site. The samples were collected near the southeastern corner of the site in order to assess the impact of groundwater seeps along the bank on the surface water quality. The location and elevation along the drainage channel where surface water samples were taken were marked with a construction stake for future reference by the surveyor.

Prior to starting the sampling procedure, the lengths of the drainage channels were inspected for evidence of surface water contamination, such as sheens on the water surface, seepage fronts along the upgradient banks of the channels, stained soils, and odors. No such evidence was found.

2.3.2 Sample Collection

Surface water samples were collected using a peristaltic pump. The tubing leading to the pump was placed beneath the water surface and kept below the water surface during sample collection to minimize aeration of the sample. Sample containers were filled directly from the pump discharge tube. The order of sample collection was from downstream to minimize the potential effects of sample collection disturbances on subsequent samples.

2.4 SAMPLE HANDLING FOR LABORATORY ANALYSIS

This section describes the handling of samples from the time of sampling until the samples were delivered to the fixed-base laboratory. For this study Curtis and Tompkins of Berkeley. California and EAL of Wheatridge, Colorado provided analytical laboratory support. Fixed-base analytical methods and field methods used for surface water samples consisted of a subset of those used for groundwater samples and are listed in Table 2.1.

2.4.1 Sample Preservation

All necessary sample containers were provided by the laboratories, and necessary chemical preservatives were added to the sample containers by the laboratories prior to shipment to the site. Samples were prepared for transportation to the laboratories by placing the samples in a cooler containing ice to maintain a shipping temperature of approximately 4 degrees centigrade (°C). Samples were delivered to Curtis and Tompkins and EAL via overnight courier to ensure that the temperature remained constant and that all sample holding times were met.

2.4.2 Sample Container and Labels

Sample containers and appropriate container lids were provided by the laboratories. The sample containers were filled as described in Sections 2.2.2 and 2.3.2, and the container lids were tightly closed. The sample label was firmly attached to the container side, and the following information was legibly and indelibly written on the label:

- Facility name;
- Sample identification;
- Sample type (e.g., groundwater);
- Sampling date:
- Sampling time:
- Preservatives added;
- Sample collector's initials; and
- Requested analyses.

2.4.3 Sample Shipment

After the samples were sealed and labeled, they were packaged for transport to the laboratories. The following packaging and labeling procedures were followed:

- Samples were packaged to prevent leakage or vaporization from their containers;
- Samples were cushioned to avoid breakage; and
- Ice was added to the shipping container to keep samples cool.

The packaged samples were delivered by overnight courier to the analytical laboratories.

2.4.4 Chain-of-Custody Control

After the samples were collected, chain-of-custody procedures were followed to establish a written record of sample handling and movement between the sampling site and the analytical laboratory. Each shipping container contained the chain-of-custody form completed by the sampling personnel. This form is a part of the permanent record for the samples and was returned with the sample analytical results. The chain-of-custody form contains the following information:

- Sample identification number;
- Sample collectors' printed names and signatures;
- Date and time of collection;
- Place and address of collection;
- Sample matrix;
- Chemical preservatives added;
- Analyses requested;
- Signatures of individuals involved in the chain of possession; and
- Inclusive dates of possession.

The chain-of-custody documentation was placed inside the shipping container so that it was immediately apparent to the laboratory personnel receiving the container without risk of being lost or damaged during transport. The shipping container was sealed so that it was obvious if the seal had been tampered with or broken.

2.5 AQUIFER TESTING

Slug tests were conducted on the two existing MWs at the site (MW-1 and MW-2). Slug tests are single-well hydraulic tests used to determine the hydraulic conductivity of an aquifer in the immediate vicinity of the tested well. This information is required to accurately estimate the velocity of groundwater and contaminants in the shallow saturated zone. Slug tests can be used for both confined and unconfined aquifers that have a transmissivity of less than 7,000 square feet per day (ft²/day).

Slug testing can be performed using either a rising head or a falling head test. At this site, two rising head and two falling head tests were performed at each well. Detailed slug testing procedures are presented in the Technical Protocol for Implementing Intrinsic Remediation with Long-Term Monitoring for Natural Attenuation of Fuel Contamination Dissolved in Ground Water (Wiedemeier et al., 1995), hereafter referred to as the Technical Protocol document.

Data obtained during slug testing were analyzed using AQTESOLV software (Geraghty & Miller, 1991) and the methods of Bouwer and Rice (1976) and Cooper *et al.* (1967) for confined conditions. The results of slug testing are presented in Section 3.3.2.2.

2.6 SURVEYING

All MPs, MWs, and surface water sampling locations were surveyed by Phillippi Engineering, Inc. (PEI) of Vacaville, California. All horizontal coordinates and ground surface elevations were surveyed to at least the nearest 0.01 foot relative to a Base datum. In addition, the top-of-casing elevation (measurement datum) for MPs was surveyed to the nearest 0.01 foot.

SECTION 3

PHYSICAL CHARACTERISTICS OF THE STUDY AREA

This section incorporates data collected by Parsons ES during field activities in support of remediation by natural attenuation (RNA) with data documented in previous reports on Beale AFB. Descriptions of regional topography, surface water hydrology, geology, and hydrogeology were taken from previous reports (CH2M Hill, 1991; LAW, 1995). Investigative techniques used to determine the physical characteristics of the site are discussed in Section 2.

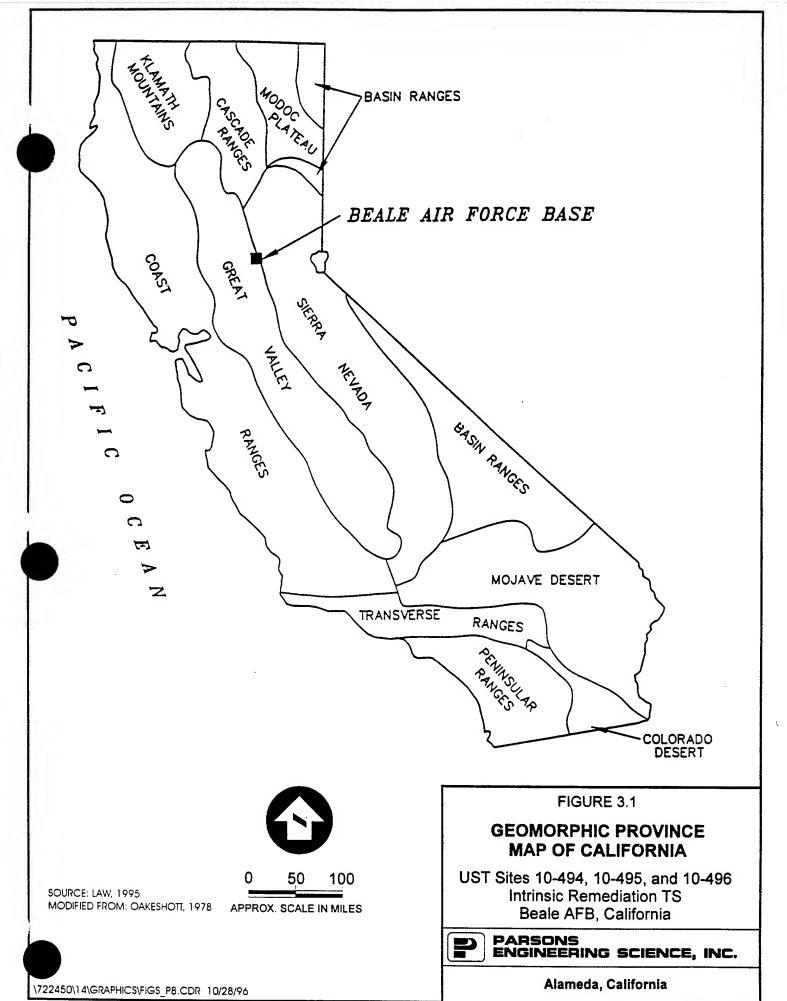
3.1 SURFACE FEATURES

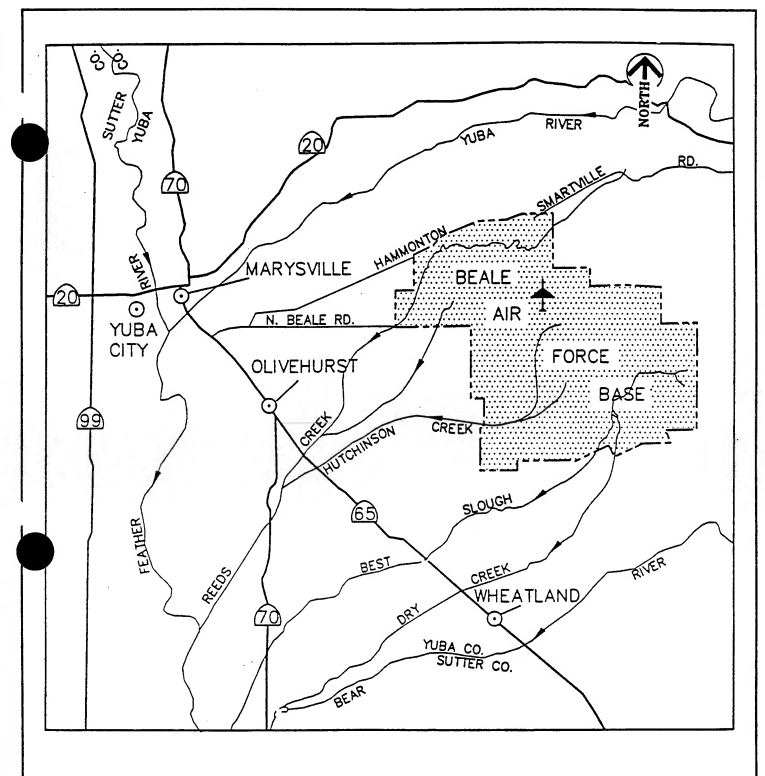
3.1.1 Topography and Surface Water Hydrology

Beale AFB is located on the eastern edge of the Sacramento Valley along the border of two geomorphic provinces: the Great (or Central) Valley to the west, and the Sierra Nevada to the east (Figure 3.1). The Great Valley Province is a nearly flat alluvial plain, approximately 450 miles long and 50 miles wide. The western portion of Beale AFB consists of generally flat plain, typical of the Great Valley Province. The Sierra Nevada Province consists of a north-northwesterly trending mountain range, approximately 400 miles long and 50 to 80 miles wide, with a shallow western slope and a steep eastern escarpment. The eastern portion of the Base has generally west-facing slopes and consists of rolling hills typical of the western foothills of the Sierra Nevada Mountains. Elevations on the Base range from approximately 80 feet above mean sea level (msl) in the west to approximately 600 feet above msl in the east.

The regional drainage pattern follows the topography, with surface water east of the Base generally flowing east to west, and surface water west of the Base generally flowing north to south. The primary drainages are the Yuba River, Bear River, Feather River, and the Sacramento River. The Yuba River, north of the Base, and the Bear River, south of the Base, flow into the Feather River, west of the Base (Figure 3.2). The Feather River drains predominately north to south and flows into the Sacramento River, which eventually flows westward through the Carquinez Straits, San Pablo Bay, and San Francisco Bay to the Pacific Ocean.

Surface drainage at Beale AFB follows the general northeast to southwest slope of the ground surface. Figure 3.3 shows the distribution of lakes, creeks, and tributaries on the Base. Much of the surface water runoff is conveyed by three main creeks: Dry Creek. Hutchison Creek, and Reeds Creek. Because of the presence of locally impermeable clay substrates in the soil profile, ephemeral vernal pools form over much of Beale AFB







SOURCE: LAW, 1995 MODIFIED FROM: PAGE, 1980 0 1.5 3
APPROX. SCALE IN MILES

FIGURE 3.2

REGIONAL SURFACE HYDROLOGY

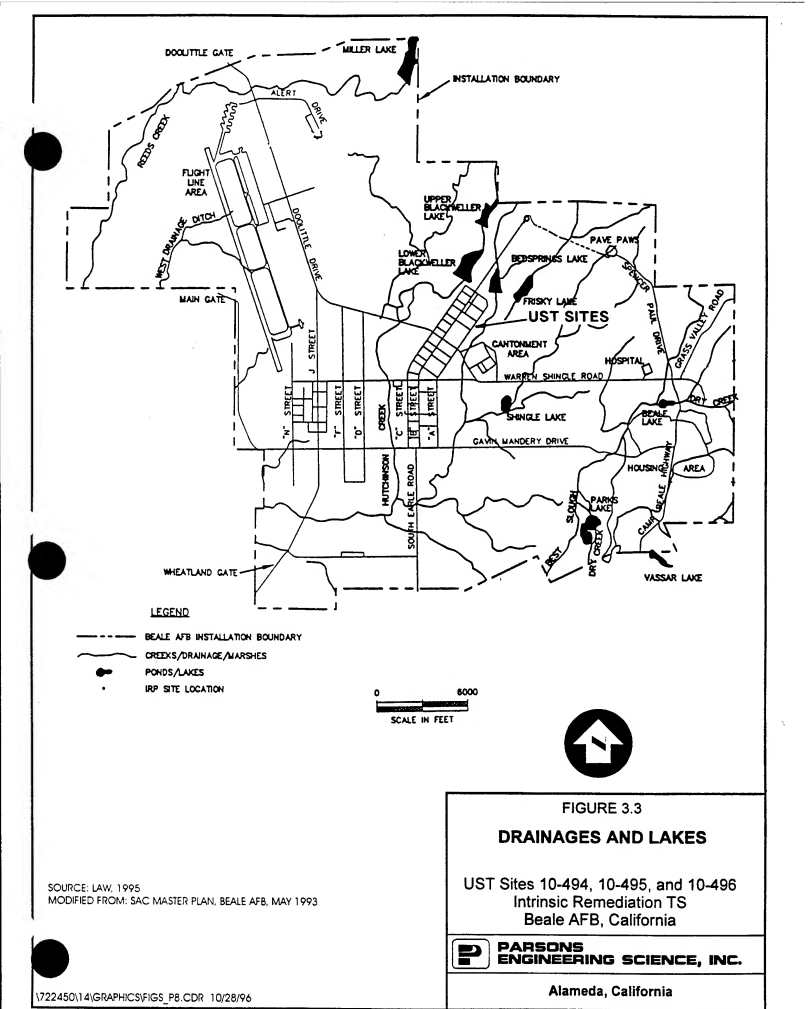
UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California



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Alameda, California





following winter rains. These pools, and most of the creeks and tributaries, dry up in the summer. Reeds Creek and Hutchison Creek join and flow into the Feather River. Dry Creek flows into the Bear River, which then flows into the Feather River.

3.1.2 Manmade Features

The land surface at the site is relatively undisturbed and is vegetated with grass. A large drainage channel, approximately 8 feet in depth and 10 to 15 feet wide, borders the site to the southeast (Figure 2.1). Several shallower drainage ditches, approximately 4 feet in depth and 5 feet wide, cut northwest to southeast and join the larger drainage channel. The smaller drainage ditches dry up in the spring after the end of the winter rains and, therefore, are assumed to exhibit losing stream features with respect to the aquifer. The large drainage channel flows year round and, therefore, is assumed to exhibit gaining stream features with respect to the aquifer. However, water depth, flow volume, and velocity are significantly lower during the summer months.

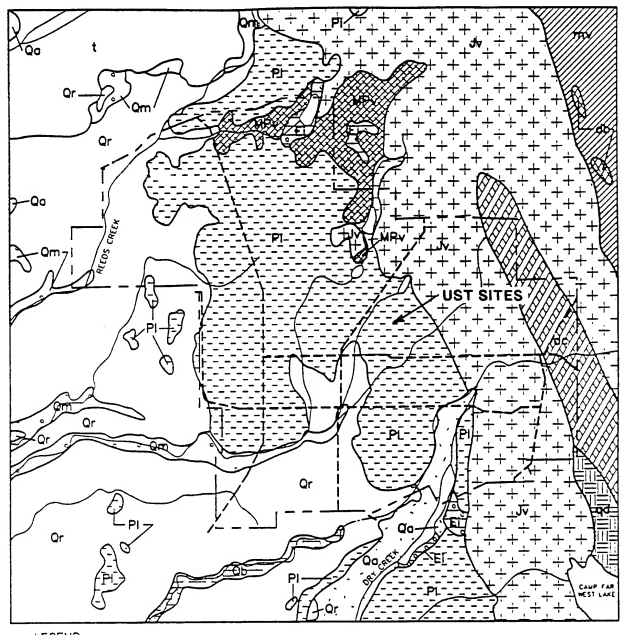
As discussed in Section 1.2, a compost pit existed at the site in the former tank excavation pit until November 1995. The fuel pipelines that supplied the USTs are inactive but are still in place. These lines run through the site parallel to the large drainage channel (Figure 2.1). The location of the pipeline is noticeable in some areas of the site due to the raised land surface.

3.2 REGIONAL GEOLOGY AND HYDROGEOLOGY

Rocks that underlie the Sacramento Valley and border the foothills of the Sierra Nevada include weakly metamorphosed plutonic, sedimentary, and volcanic rocks of Paleozoic and Mesozoic age; sedimentary and volcanic rocks of Mesozoic and Cenozoic age; and unconsolidated alluvium of Quaternary age.

The geology of the area reflects the boundary between the Sierra Nevada and the Great Valley geomorphic provinces. The Sierra Nevada Province is composed of a westward tilting crustal block of the Sierra batholith consisting of primarily Jurassic to Cretaceous plutonic rocks. To the west of these plutonic rocks is a belt of weakly metamorphosed sedimentary and volcanic rocks of Paleozoic and Mesozoic age. The Great Valley Province is composed of an elongated structural trough filled with a thick sequence of Jurassic to Recent age sediments. In the eastern part of the valley, including Beale AFB, these deposits are derived from and overlie the metamorphic and igneous rocks of the Sierra Nevada. The sediment deposits reach a maximum thickness of 60,000 feet in the Central Valley.

The geologic units in the vicinity of the Base include undifferentiated Cretaceous units and Tertiary and Quaternary formations, as shown on Figure 3.4. The Tertiary and Quaternary units found in the vicinity of the Base, from oldest to youngest, include the Capay, Ione, Wheatland, Mehrten, Neroly, Laguna, Riverbank, and Modesto Formations, and Holocene deposits. A description of these units is included in the generalized stratigraphic column presented in Table 3.1.



LEGEND

t Holocene Dredge or Mine Tailings

Qa

Qm

Holocene Natural Levee and Channel Deposits

Holocene Basin Deposits

Pleistocene Modesto Formation

Qr Pleistocene Riverbank Formation

Pliocene Laguna Formation

Miocene-Pliocene
Volcanic Rocks

Eocene Ione Formation

Jurassic Volcanic Rocks Predominantly Metamorphosed Mafic to Felsic Pyroclastic Rock

Jurassic Volcanic Rocks Predominantly Pyroclastic and Volcanic Rock

Jurassic Quartz Diorite

Jurassic Dike Complex

> Jurassic Massive Diabase

--- Road

SOURCE: LAW, 1995 MODIFIED FROM: SAUCEDO AND WAGNER, 1992

FIGURE 3.4

BASEWIDE GEOLOGIC MAP

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California



PARSONS ENGINEERING SCIENCE, INC.

Alameda, California

10/28/96 \tag{722450\14\GRAPHICS\FIGS_P8.CDR}

APPROXIMATE SCALE IN FEET

TABLE 3.1 GENERALIZED STRATIGRAPHIC COLUMN

UST Sites 10-494, 10-495, and 10-496

Intrinsic Remediation TS Beale AFB, California

Formation	Age of Formation	Description	References
Recent Deposit	0 to 11,000 years Holocene	Unconsolidated alluvial silt, sand, and gravel and overbank clay, silt, and fine-grained sand.	Page, 1980
Modesto	9,000 to 42,000 years Holocene to Pleistocene	Alluvial, fluvial, and possibly lacustrine sand, silt, and clay with minor gravel	Busacca, 1982
Riverbank	150,000 to 450,000 years Pleistocene	Fluvial silt, sand, and gravel derived from erosion of the Laguna Formation	Busacca, 1982
Laguna	1.6 to 3.4 m.y. Pleistocene to Pliocene	Heterogeneous assemblage of silt, clay, sand and minor siliceous gravel beds. Predominantly reddish to yellowish-brown silt to sand silt with abundant clay and minor intervals of tuff.	Redwine, 1972
Neroly	5 to 22 m.y. Pliocene to Late Miocene	Volcanic fluvial deposits ranging from volcanoclastic cobble conglomerate to sandstone and clayey siltstone, with minor intervals of tuff.	Unruh, 1990
Mehrten	5 to 22 m.y. Pliocene to Late Miocene	Volcanic mudflow debris and pyroclastic flows deposited on the western front of the Sierra Nevada. The sediments are predominantly andesitic, but range from basaltic to rhyolitic.	Clark & Anderson, 1938; Redwine, 1972
Wheatland	25 to 38 m.y. Oligocene	Predominantly a tuffaceous sandstone with basal conglomerate containing marine fossils. The sandstone is overlain by thinly bedded fissile, tuffaceous lignitic shale. Near the top of the section lies a pink volcanic mudflow with subangular volcanic clasts. The top of the formation consists of well-bedded spheroidal shale with interbeds of lignite containing carbonized reeds.	Unruh, 1990
Ione	44 to 45 m.y. Early to Middle Eocene	A light-colored anauzitic sandstone and clay- stone, dark reddish-brown or brown ferruginous sandstone, quartz, and metamorphic gravel, and conglomerate with minor lignite.	Redwine, 1972; Unruh, 1990
Capay	50 to 55 m.y. Early Eocene	Dark greenish-gray, fossiliferous siltstone, clayey siltstone, and sandy siltstone with occasional sand and fine gravel.	
Notes:			Source: Law 1995

Surface soils consist primarily of three different types of loam: moderately deep to deep loam formed in alluvium (Perkins loam); shallow to moderately deep gravelly loam with cobbles (Pardee loam); and small areas of shallow to moderately deep fine-grained soils (Auburn loam). A map of surface soils is shown on Figure 3.5. Permeability is typically low to moderate.

Water-bearing deposits in the Sacramento Valley and in the vicinity of the Base consist primarily of the non-marine, valley-filling late Tertiary and Quaternary sediments. Water-bearing deposits in the vicinity of Beale AFB are the Neroly, Laguna, Riverbank, and Modesto Formations, and Holocene-age alluvial deposits. These water-bearing zones range in thickness from approximately 1,000 feet in the vicinity of the Feather River to less than 300 feet in the vicinity of the Sierra Nevada foothills. At Beale AFB, the volcaniclastic Neroly sediments beneath the Laguna Formation are the most productive water-bearing zones. Groundwater also is found in the metamorphosed volcanic rocks of the Sierra Nevada foothills, primarily in the fractures.

In the Sacramento Valley and on the Base, groundwater conditions are generally unconfined. However, locally confined and/or perched zones can occur in areas where sands and gravels are found with less permeable silts and clays. In general, flow in the Sacramento Valley is toward the center of the valley. However, groundwater pumping has depressed the groundwater table in portions of the Sacramento Valley and has created localized flow conditions that do not follow the regional topography.

Groundwater elevations at the Base range from approximately 20 feet above msl in the western area of the Base to approximately 140 feet above msl in the eastern area (Figure 3.6). Groundwater flow is generally to the southwest. Extensive pumping in the area has influenced groundwater flow directions.

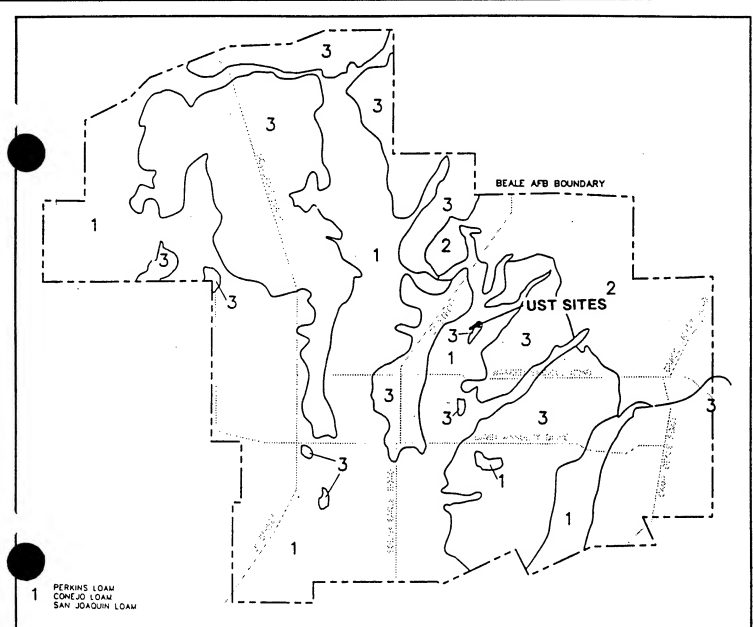
3.3 SITE GEOLOGY AND HYDROGEOLOGY

3.3.1 Lithology and Stratigraphic Relationships

The uppermost geologic unit in the western portion of Beale AFB, which includes the UST Site, has been mapped as Laguna Formation. This formation consists of a Plio-Pleistocence alluvial sequence of silt, sand, clay, and unsorted gravel. These deposits overlie older volcanic rocks that outcrop in the Sierra Nevada foothills to the east.

Evaluation of the site-specific geology is based on the logs from soil borings advanced during the current investigation and during a preliminary investigation by Metcalf & Eddy (1995) (Figures 3.7 through 3.9 and Appendix A).

Soils in the vicinity of the UST Site consist of interbedded clays, silts, sands, and gravels. The cross-sections (Figures 3.8 and 3.9) generally show a surficial layer across the site consisting of gravelly and sandy clay and silt to a depth of approximately 5 feet bgs. This layer is underlain by a unit of coarse-grained sediments, consisting of silty and/or gravelly fine- to coarse-grained sands. The gravels and sands generally end at a depth of 8 to 10 feet bgs and are underlain by silty clay or clayey silt to approximately 11 to 15 feet bgs. A lower



2 AUBURN LOAM ARGONAUT-AUBURN COMPLEX AUBURN-SOBRANTE COMPLEX

3 PARDEE LOAM
PARDEE VARIANT COMPLEX
REDDING-CORNING COMPLEX
REDDING LOAM





FIGURE 3.5

SURFACE SOILS

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

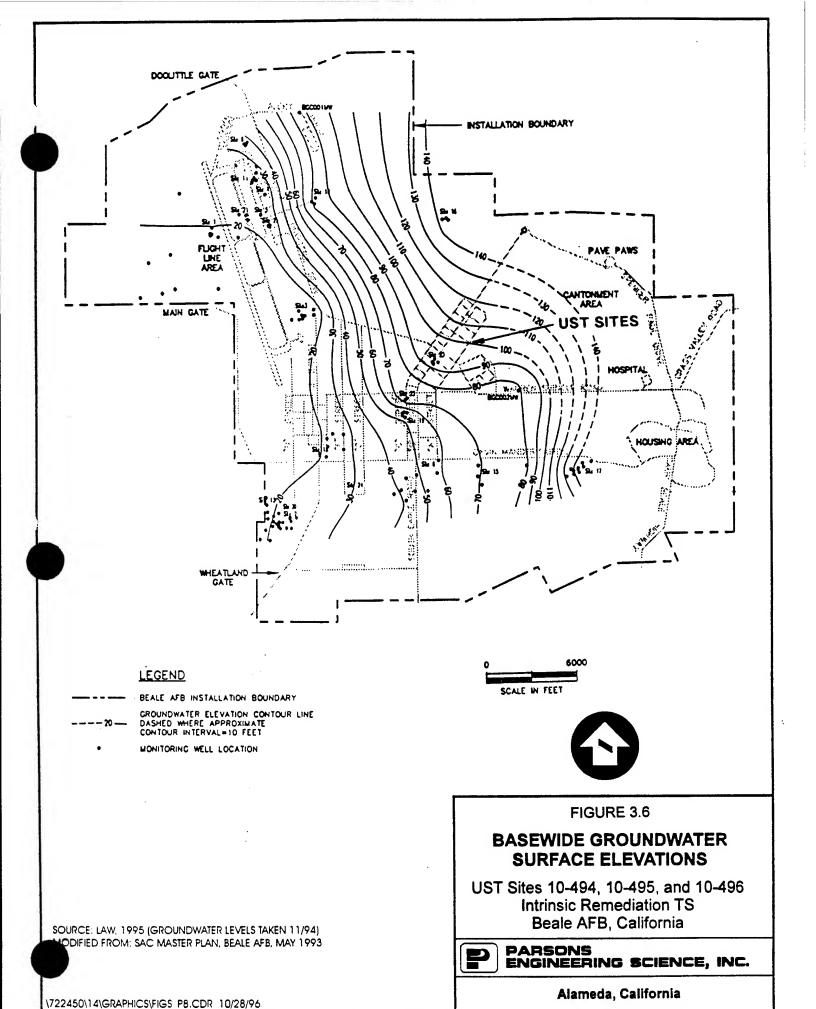


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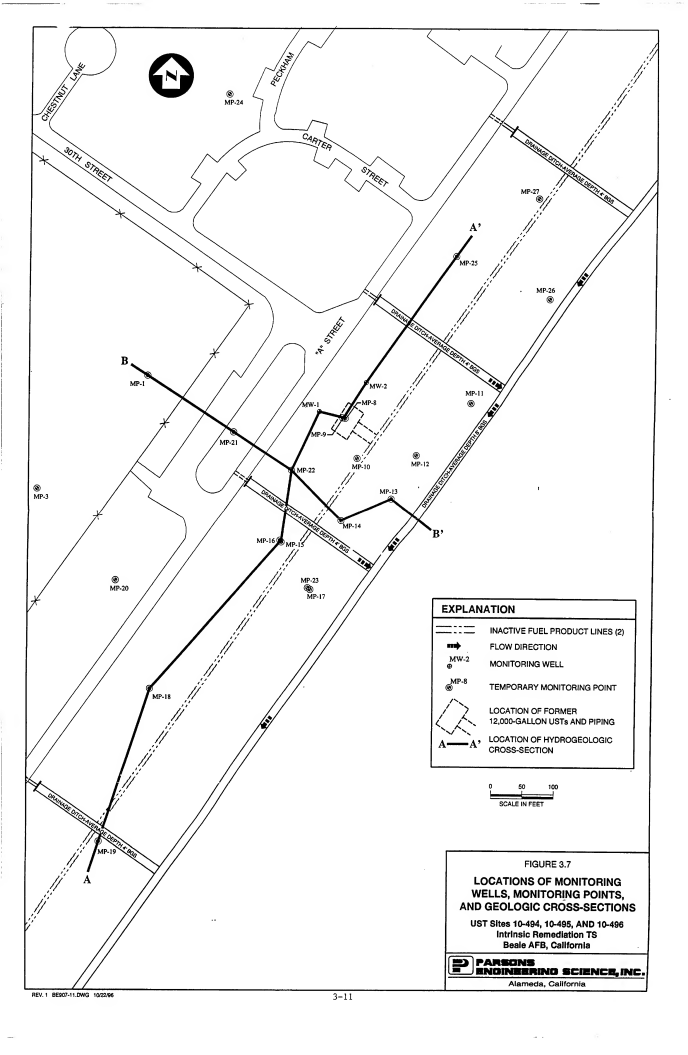
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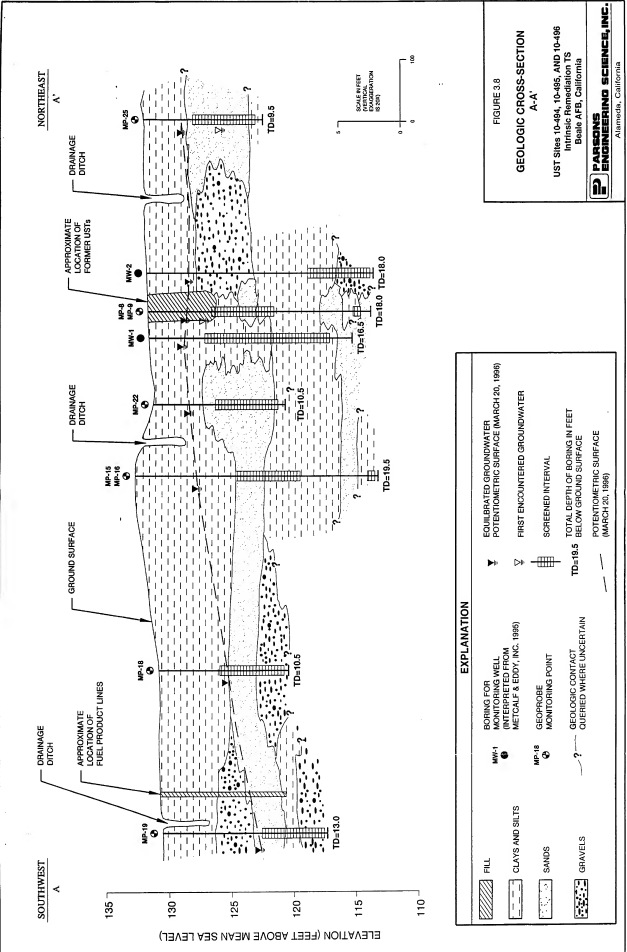
SOURCE: LAW, 1995 MODIFIED FROM: USDA, SOIL CONSERVATION SERVICE, 1985

\722450\14\GRAPHICS\FIGS_P8.CDR 10/28/96



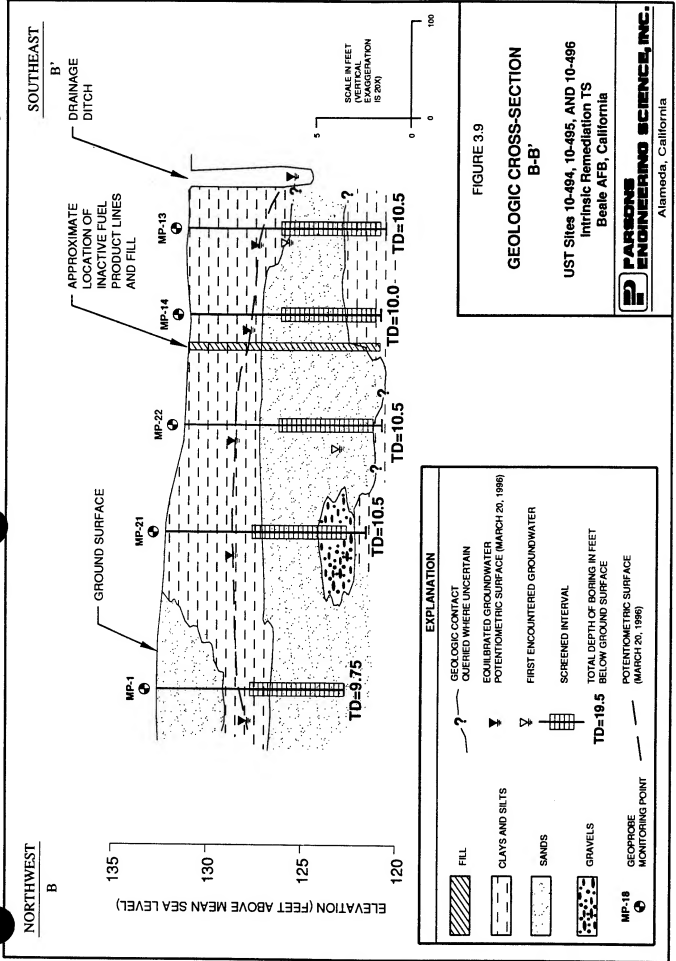
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layer of sands and gravels was encountered beneath the layer of silty clay and clayey silt in some boreholes, but it is not known if this lower layer of sand and gravel is laterally continuous.

Soil samples collected and analyzed for grain-size distribution were consistent with the field geologist's interpretations. Soil samples were collected for grain-size distribution analysis from both the surficial gravelly and sandy clay and silt (at MP-24 at 3.0 feet bgs) and the underlying saturated coarse-grained unit (at MP-25 at 7.0 feet bgs). The following results were obtained: 5.5 % gravel, 31.5% sand, 6.0% silt, and 57.0% clay at MP-24; and 24.6% gravel, 59.8% sand, 1.0% silt, and 14.6% clay at MP-25.

Investigations indicate that the shallow soil units are generally flat-lying and laterally continuous across the site. The surface layer of sandy and gravelly silts and clays appears to act locally as an aquitard, confining the upper coarse-grained layer. The layer of silty clay to clayey silt underlying the coarse-grained sediments also may act as an aquitard, separating the upper and lower coarse-grained units.

At the southern end of the site, near MP-19, the upper and lower coarse-grained units may be connected. The coarse-grained units encountered in MP-19 are separated by a 1-foot thick layer of sandy, gravelly clay. However, the coarse-grained materials encountered in MP-19 may also be a thick section of the upper coarse-grained unit, with the silty clay to clayey silt layer underlying the coarse-grained unit at a depth greater than the depth of the borehole. The aquifer also appears to become unconfined at the southern end of the site near MP-18 and MP-19.

3.3.2 Groundwater Hydraulics

3.3.2.1 Flow Direction and Gradient

The groundwater levels at the site varied from 123.43 to 131.26 feet above msl in March 1996, and 123.01 to 129.61 feet above msl in May 1996 (Table 3.2). These levels are about 20 to 30 feet higher than expected based on the regional groundwater potentiometric surface map (Figure 3.6). The shallow, coarse-grained, water-bearing unit encountered at the site may be a "perched" aquifer or the regional groundwater potentiometric surface map may simply be inaccurate near the UST Site because it is based on sparse data in the northeastern portion of the Base.

Groundwater is generally encountered first in the upper coarse-grained unit or at the unit's contact with the overlying silt and clay layer (Figures 3.8 and 3.9), but often rises to an equilibrated level at or above the contact between the upper coarse-grained layer and the overlying silt and clay. Groundwater levels measured in the MWs and MPs at the site were between 2.15 and 6.97 feet bgs in March 1996, and between 3.87 to 7.39 feet bgs in May 1996. A summary of historical groundwater measurements is presented in Table 3.2. A groundwater elevation/potentiometric surface map for the site for March 1996 is presented on Figure 3.10. This map is based on groundwater levels measured in MW-1 and the shallow MPs, and on the surface water levels measured in the large drainage channel in March 1996. Because the MWs at the site were installed in February 1995 and have not been sampled or

TABLE 3.2 SUMMARY OF HISTORICAL WATER LEVEL MEASUREMENTS

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

				Top of		Danahaa	Groundwater/Surface
Location	Date	Northing	Easting	Casing Elevation	Ground	Depth to Groundwater	Water Potentiometric Surface
Location	Date	(ft)	(ft)	(ft msl)	(ft msl)	(ft btoc) ²	(ft msl) ¹
MP-01	Mar-96	530590.0	2174967.0		132.40	4.54	128.09
	May-96					5.01	127.62
MP-03	Mar-96 May-96	530403.4	2174793.8	131.43	130.20	3.38 5.10	128.05 126.33
MP-08	Mar-96	530531.9	2175281.3	n.a.	131.80	dry	•
MP-09	Mar-96 May-96	530531.4	2175282.2	132.01	131.80	3.34 4.59	128.67 127.42
MP-10	Mar-96 May-96	530464.8	2175302.9	132.09	131.20	3.95 5.23	128.14 126.86
MP-11	Mar-96 May-96	530558.0	2175483.4	133.52	132.40	5.50 6.12	128.02 127.40
MP-12	Mar-96 May-96	. 530472.9	2175398.2	131.67	131.40	4.62 5.32	127.05 126.35
MP-13	Mar-96 May-96	530399.1	2175356.1	131.20	131.10	3.98 4.81	127.22 126.39
MP-14	Mar-96 May-96	530360.9	2175277.2	131.38	130.90	3.83 5.12	127.55 126.26
MP-15	Mar-96 May-96	530324.5	2175181.7	134.91	132.90	7.19 8.81	127.72 126.10
MP-16	Mar-96	530326.5	2175179.4	n.a.	132.90	dry	-
MP-17	Mar-96 May-96	530248.5	2175227.7	132.54	131.90	6.73 7.30	125.81 125.24
MP-18	Mar-96 May-96	530086.1	2174977.7	131.15	130.90	5.36 6.38	125.79 124.77
MP-19	Mar-96 May-96	529831.4	2174896.5	132.80	130.40	9.37 9.79	123.43 123.01
MP-20	Mar-96 May-96	530258.1	2174918.9	130.89	130.30	4.12 5.45	126.77 125.44
MP-21	Mar-96 May-96	530502.8	2175103.4	132.71	132.20	3.95 5.70	128.76 127.01
MP-22	Mar-96 May-96	530442.8	2175195.6	131.39	131.20	2.87 4.47	128.52 126.92
MP-23	Mar-96	530250.1	2175225.9	n.a.	131.90	dry	-

TABLE 3.2 (continued) SUMMARY OF HISTORICAL WATER LEVEL MEASUREMENTS

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

				Top of Casing		Depth to	Groundwater/Surface Water Potentiometric
Location	Date	Northing	Easting	Elevation	Ground	Groundwater	Surface
		(ft)	(ft)	(ft msl) ¹	(ft msl)	(ft btoc) ²	(ft msl) ¹
MP-24	Mar-96 May-96	531049.9	2175090.2	134.92	134.30	3.66 5.91	131.26 129.01
MP-25	Mar-96 May-96	530793.0	2175457.6	133.12	132.20	3.86 4.85	129.26 128.27
MP-26	Mar-96 May-96	530727.5	2175607.0	132.86	131.80	4.88 5.24	127.98 127.62
MP-27	Mar-96 May-96	530886.9	2175596.8	133.10	132.90	3.81 4.49	129.29 128.61
MW-01	Feb-95 Mar-96 May-96	530539.3	2175241.9	131.65	131.80	5.35 2.42 4.12	126.30 129.23 127.53
MW-02	Feb-95 Mar-96 May-96	530588.3	2175316.3	131.81	131.90	5.31 3.36 4.86	126.50 128.45 126.95
SW-01	Mar-96 May-96	530265.1	2175312.8	n.a.	124.90	n.a.	124.90 124.57 (est.)
SW-02	Mar-96 May-96	530315.0	2175294.0	n.a.	127.30	n.a.	127.30 126.97 (est.)
SW-03	Mar-96 May-96	530308.0	2175344.0	n.a.	125.00	n.a.	125.00 124.67 (est.)

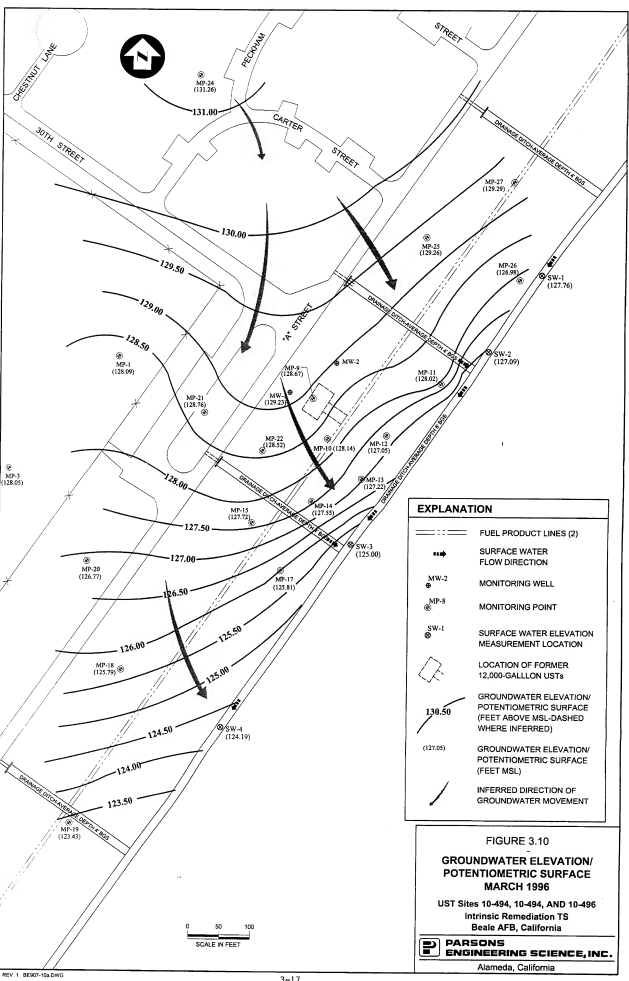
Notes

¹ ft msl - feet above mean sea level.

² ft btoc - feet below top of casing.

n.a. - not applicable

est. - estimated value



3-17

monitored regularly since then, the magnitude of the seasonal variations in the groundwater levels at the site is not known.

The average gradient calculated from water levels in March 1996 was approximately 0.0097 foot per foot (ft/ft) toward the south-southeast in the vicinity of the site. The steepest gradient at the site, 0.029 ft/ft, was found in the vicinity of MP-14. A slight variation was noted in the groundwater flow direction and gradient between March and May 1996, with the flow taking a slightly more southerly direction, and the average gradient dropping to 0.0070 ft/ft. Somewhat higher variations are expected to exist between the wettest winter months and the driest summer months. Additional groundwater level measurements collected throughout the year would be required to determine seasonal variations in groundwater flow direction and gradient.

The groundwater flow direction and gradient appear to be significantly affected by the large drainage channel, which contains water year round and likely intersects the upper coarse-grained layer. Groundwater is assumed to discharge to this channel, based on the groundwater levels measured in the surrounding MPs and the surface water level measured in the channel. This interpretation is consistent with a general analysis by Strack (1989). who found that if a drainage ditch penetrates approximately 10 percent of an aquifer thickness, over 94 percent of the groundwater discharges into the ditch.

The shallower drainage ditches northeast and southwest of the site do not appear to intersect the upper water-bearing unit; however, water in these ditches may contribute to recharge of the unit. In March 1996, water was measured at 6 to 8 inches deep in the large channel and was flowing at a rate of approximately 1 foot per second (ft/s). In the shallow ditch southwest of the site, stagnant water was measured at 1 to 3 inches deep. No water was observed in the shallow northwest ditch.

It does not appear that the upper and lower coarse-grained units are hydrogeologically connected across most of the site. The groundwater level in MW-2 in May 1996 was lower than groundwater levels measured in MW-1, MP-9, and MP-11, yet the regional potentiometric surface map indicates that MW-2 would be upgradient or slightly cross-gradient from these locations if they were hydrogeologically connected. In addition, the deep MPs screened within the intervening clayey silt did not produce any groundwater, suggesting this layer acts as an aquitard or at least significantly restricts vertical groundwater movement. However, as discussed in Section 3.3.1, the two units may be connected at the southern end of the site near MP-19.

3.3.2.2 Hydraulic Conductivity

Parsons ES estimated the hydraulic conductivity at wells MW-1 and MW-2 using both rising and falling head slug tests and the straight-line method of Bouwer and Rice (1976) and type-curve matching method of Cooper *et al.* (1967). The Bouwer and Rice method, although generally used for unconfined aquifers, can be used for confined conditions with limited accuracy (Bouwer, 1989).

The results of these slug tests are summarized in Table 3.3 and detailed in Appendix D. Input files and output files from the slug tests are provided in Appendix E. The average hydraulic conductivity obtained in rising head tests at MW-1, fully penetrating the upper water bearing unit, was 8.8 x 10⁻⁵ foot per second (ft/s) (2.7 x 10⁻³ centimeter per second [cm/s]). The average hydraulic conductivity obtained in rising and falling head tests at MW-2, partially penetrating the lower coarse-grained unit, was 6.7 x 10⁻⁵ ft/s (2.0 x 10⁻³ cm/s). These values are consistent with typical conductivities for silty sands to clean sands. The average hydraulic conductivity from the rising head tests conducted at MW-1 was used as the initial average hydraulic conductivity for the site in the groundwater model presented in Section 5.

The falling head test at MW-1 was not analyzed because of difficulties during the test. The slug was not in place at a discrete time, and water spilled over the top of the casing during placement of the slug.

3.3.2.3 Effective Porosity

Because of the difficulty involved in accurately determining effective porosity, accepted literature values for the type of soil comprising the shallow saturated zone were used. Walton (1988) give ranges of effective porosity for gravelly and coarse sands of 0.20 to 0.35. Freeze and Cherry (1979) report ranges of effective porosities for sand of 0.25 to 0.50. Because the presence of silt tends to decrease the effective porosity, and because lower effective porosities result in higher computed advective groundwater velocities, an effective porosity of 0.25 was assumed for this project.

3.3.2.4 Advective Groundwater Velocity

The advective velocity of groundwater in the direction parallel to groundwater flow is given by:

$$\overline{v} = \frac{K}{n_e} \frac{dH}{dL}$$

Where: v = Average advective groundwater velocity (seepage velocity)

K = Hydraulic conductivity

dH/dL = Gradient

 $n_{\rm e}$ = Effective porosity

Using this relationship in conjunction with site-specific data, the advective groundwater velocity at the site in March 1996 was estimated at approximately 80 to 120 feet per year (ft/year). Advective groundwater velocities are likely to be lower during the drier summer months, when the gradient would be expected to be lower.

3.3.2.5 Preferential Flow Paths

The large drainage channel appears to exert significant influence on groundwater flow at the site. In addition, the shape of the groundwater potentiometric surface contours indicates that a more transmissive zone extends from the northwest (near MP-24), toward the former UST

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TABLE 3.3 HYDRAULIC TESTING RESULTS MARCH 1996

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

Location	Test Type	Analysis Method	Hydraulic Conductivity (ft/sec)
MW-1	Slug Test, Rising Head #1	Cooper-Bredehoeft- Papadopulos	7.84E-05
·	Slug Test, Rising Head #2	Cooper-Bredehoeft- Papadopulos	9.75E-05
MW-2	Slug Test, Falling Head #1	Bouwer and Rice	9.09E-05
	Slug Test, Rising Head #1	Bouwer and Rice	5.79E-05
	Slug Test, Falling Head #2	Bouwer and Rice	6.11E-05
	Slug Test, Rising Head #2	Bouwer and Rice	5.71E-05

location (near MW-1), and then southwest of the site (toward MP-15). This interpretation was supported by the model calibration results discussed in Section 5.

In addition, a filled excavation for the subsurface fuel pipelines that cross the site from northeast to southwest may potentially create a preferential flow path not observable on the sitewide groundwater potentiometric surface map. The depth of the back-filled excavation in the site vicinity is unknown, but could be as deep as 12 feet bgs. The nature of the fill is also unknown.

3.3.2 Groundwater Use

The closest wells used for domestic, industrial, and irrigation purposes are located 3 miles west of the site along the western boundary of the Base (CH2M Hill, 1991). These wells are screened in water-bearing zones between 125 to 360 feet bgs.

3.4 CLIMATE

Beale AFB experiences an interior Mediterranean climate characterized by hot, dry summers and cool, wet winters. Temperature extremes range from a low of 20 degrees Fahrenheit (°F) to a high of 113 °F. The Base receives approximately 95 percent of its annual precipitation between October and April. Total mean annual precipitation is 22.2 inches per year.

The annual evapotranspiration (ET) rate, 66.5 inches per year, coupled with the yearly precipitation yields a net precipitation of minus 44.3 inches per year. This implies that precipitation has little chance to percolate to regional groundwater. However, approximately 95 percent of the total rainfall occurs during the winter and spring months when ET rates are relatively low, so some recharge probably occurs on a seasonal basis.

SECTION 4

NATURE AND EXTENT OF CONTAMINATION AND SOIL, GROUNDWATER, AND SURFACE WATER CHEMISTRY

4.1 SOURCES OF CONTAMINATION

The primary sources of contamination at the UST Site were fuel leaks from the three former gasoline USTs. Although no spills or leaks have been documented, site investigations have identified fuel-related contamination in the soil and groundwater near the former UST location. All three tanks were reported to have been removed prior to 1962.

The excavation pit was left open after tank removal activities and was used as a compost pile for landscape waste until November 1995. In November 1995, the compost pile was removed, the excavation pit was filled with clean soil, and the site was completely regraded.

Two 4-inch diameter pipes used to supply the USTs until the end of World War II run through the site just east of the former UST location, parallel to A Street. These fuel pipelines are inactive but still in place. Although no spills or leaks have been documented, the product lines or distribution piping from the lines to the USTs could have been a source of contamination.

In December 1994, investigations by LAW (1995) identified hydrocarbon contamination at soil vapor points (SVPs) installed near the former USTs. In February 1995, soil and groundwater investigations by Metcalf & Eddy (1995) also identified hydrocarbon contamination in soil boring and test pit samples located around the excavation pit.

Two monitoring wells, MW-1 and MW-2, were installed near the excavation pit. Although hydrocarbon contamination was not detected in groundwater samples from these wells, groundwater samples from three test pits (TP-8, TP-9, and TP-12) and one soil boring (SB-3) showed hydrocarbon contamination.

A more detailed description of past investigations is outlined in the work plan for the TS (Parsons ES, 1995). A summary of the soil and groundwater sampling results from prior investigations and this investigation is presented in the following sections. Analytical results from previous investigations are summarized in Appendix B.

4.2 SOURCE AND SOIL CHEMISTRY

4.2.1 Mobile LNAPL Contamination

Mobile LNAPL is defined as LNAPL that is free to flow in the aquifer and that will flow from the aquifer matrix into a well under the influence of gravity. Mobile LNAPL may be present in the shallow groundwater (3 to 6 feet bgs), as evidenced by visual indications of free product during drilling of SB-6 and soil vapor points 132 and 142 in February 1995 (Metcalf & Eddy, 1995). However, no mobile LNAPL was encountered during soil. groundwater, or surface water sampling performed by Parsons ES personnel in March 1996; therefore, any mobile LNAPL at the site is likely to be very limited in extent.

In addition, the low concentrations of dissolved BTEX in groundwater at the site (discussed in Section 4.3) suggest that any mobile LNAPL that may be present is likely to be highly weathered, probably due to the age of the fuel releases.

4.2.2 Residual LNAPL Contamination

Residual LNAPL is defined as the LNAPL that is trapped in the aquifer by the processes of cohesion and capillarity, and therefore will not flow within the aquifer and will not flow from the aquifer matrix into a well under the influence of gravity. At this site, any residual LNAPL consists of fuel hydrocarbons primarily derived from gasoline.

Petroleum contamination was detected at low concentrations in several soil samples collected at the site (Section 4.2.3); however, the majority of detected soil contamination is believed to have been derived from the sorption of contaminants in the groundwater to soil because most soil samples were collected near the capillary fringe or in the saturated zone due to the high groundwater levels encountered at the site. In all cases, the residual soil contamination appeared to be highly weathered. The maximum concentrations of TPH and BTEX in soil, 780 mg/kg and 2.2 mg/kg, respectively, were measured in the sample collected at MP-10 at 5.5 feet bgs in the unsaturated zone. This sample was collected just outside the excavation pit boundary at a depth corresponding with the bottom of the former USTs. This was the only MP where field observations indicated that soil contamination was in the vadose zone, suggesting that residual contamination in the vadose zone is limited to the fringes of the excavation boundary. Previous site investigation soil and soil gas sampling results also indicated that residual soil contamination is probably limited to just outside the excavation pit boundary.

4.2.3 Soil and Soil Gas BTEX and Fuel Hydrocarbon Contamination

Soil samples were collected for lithologic description and organic and inorganic analyses at the UST Site in February 1995 (Metcalf & Eddy, 1995) and during this investigation in February/March 1996. In addition, soil vapor screening was conducted in December 1994 (LAW, 1995). The soil and soil vapor sampling analytical data from the prior investigations are included in Appendix B.

During the soil vapor survey conducted in December 1994, soil vapor samples were collected from a total of seven soil vapor points (SVPs) (132, 133, 134, and 142 through 145) at depths

between 5 and 8 feet bgs. Sampling locations are shown on Figure 2.1. Soil vapor samples were analyzed for total petroleum hydrocarbons (TPH) by EPA Method SW8015 (modified), BTEX by EPA Method SW8020, and halogenated volatile organic compounds (VOCs) by EPA Method SW8010. The maximum detected concentrations were 27,120 parts per million by volume (ppmv) TPH, 87.66 ppmv benzene, and 132.45 ppmv total BTEX, measured at SVP 132, located near MW-2 and north of the excavation pit. The highest detections from the soil vapor survey were generally in soils nearest to and downgradient from the UST excavation pit. No halogenated VOCs were detected in any soil vapor sample.

During a subsequent site investigation in February 1995, soil samples were collected from seven soil boreholes (SB-1 through SB-7) and one test pit (TP-10). Soil samples were analyzed for TPH as gasoline (TPH-g); TPH as diesel (TPH-d); BTEX; and total lead. The maximum detected concentrations were 180 milligrams per kilogram (mg/kg) TPH-g (SB-4 at 5.5 feet bgs) and 25 mg/kg TPH-d (SB-6 at 5.5 feet bgs). SB-4 and SB-6 are both located just outside and downgradient from the former excavation pit (Figure 2.1). No BTEX compounds were detected in soil samples. Total lead concentrations did not exceed the background soil concentration range of 17 to 35 mg/kg used at Beale AFB (LAW, 1995).

Eleven soil samples were collected in February/March 1996 by Parsons ES personnel during investigation activities for this study. Soil sample depths ranged from 4.0 to 9.5 feet bgs. Samples were analyzed for TPH (quantified as gasoline, diesel, jet fuel, and motor oil). BTEX, and the trimethylbenzene (TMB) isomers. Soil analytical results are summarized in Table 4.1. TPH concentrations ranged from below detection limits to a maximum of 780 mg/kg TPH-g at MP-10. The laboratory reported most TPH quantitations as not matching the calibration standard and/or heavier than the standard, probably due to the highly weathered nature of the residual contamination. BTEX concentrations ranged from below detection limits to 2.2 mg/kg (consisting entirely of xylenes) at MP-10. Benzene, toluene, and ethylbenzene were not detected in any soil samples.

4.2.4 Total Organic Carbon

TOC concentrations are used to estimate the amount of organic matter sorbed on soil particles or trapped in the interstitial passages of a soil matrix. The TOC concentration in the saturated zone is an important parameter used to estimate the amount of contaminant that could potentially be sorbed to the aquifer matrix. Sorption results in retardation of the contaminant plume relative to the average advective groundwater velocity.

During this investigation, three soil samples collected from the saturated zone were analyzed for TOC (MP-19, MP-24, and MP-25). All locations were outside the area of contamination. The TOC concentrations ranged from 90 mg/kg to 2,200 mg/kg.

4.3 GROUNDWATER AND SURFACE WATER CHEMISTRY

4.3.1 Fuel Hydrocarbon Contamination

Laboratory analytical results from sampling conducted as part of this TS investigation confirmed the presence of fuel hydrocarbon contamination dissolved in the shallow groundwater at the UST Site. Total dissolved BTEX was detected in groundwater samples

FUEL HYDROCARBON COMPOUNDS DETECTED IN SOIL SAMPLES **MARCH 1996** TABLE 4.1

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

	Analyses:	Tota	Total Petroleum Hydrocarbons	Hydrocart	suoc			BTEX			0	ther Volatil	Other Volatile Aromatics	
	Method:		USEPA SW8015M	W8015M					USEPA	USEPA SW5030/SW8020A	W8020A			
								Ethyl-	Total					Total
	Analyte:	TPH-g	TPH-JP5	P-H-L	TPH-mo	Benzene	Toluene	penzene	Xylenes	BTEX	1,2,3-TMB	1,2,3-TMB 1,2,4-TMB 1,3,5-TMB	1,3,5-TMB	TMB
	Depth													
Location	(ft bgs)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)
MP-01	5.0	<1.1	 -	4.4 YH	51	<0.0057	<0.0057	<0.0057	<0.0114	<0.029	<0.0057	<0.0057	<0.0057	<0.017
MP-09	0.9	<1.2	2.3	8.8 YH	15 YL	<0.0058	<0.0058	<0.0058	<0.0116	<0.029	<0.0058	<0.0058	<0.0058	<0.017
MP-10	5.5	HA 08L	Y 011	120 Y	9:5>	<0.110	<0.110	<0.110	2.2	2.2	<0.11	0.18	<0.11	0.18
MP-11	7.0	40 YH	5.5 YH	10 Y	<5.7	<0.014	<0.014	<0.014	0.16	0.16	0.11	0.085	0.016	0.21
MP-12	4.0	<1.2	<1.2	<1.2	9>	<0.006	>0.006	<0.006	<0.012	<0.030	<0.006	<0.006	<0.006	<0.018
MP-13	6.5	<1.2	λL	4.7 YL	<5.8	<0.0058	<0.0058	<0.0058	<0.0116	<0.029	<0.0058	<0.0058	<0.0058	<0.017
MP-14	7.0	21 Y	Y 28	33 YL	21 YL	<0.0059	<0.0059	<0.0059	0.081	0.081	0.12	0.16	0.13	0.41
MP-15	9.5	22 Y	λ ει	110 Y	<5.6	<0.0056	<0.0056	<0.0056	0.086	0.086	0.19	<0.0056	0.090	0.28
MP-19	7.0	<1.1	1.3 Z	1.3 Z	<5.5	<0.0055	<0.0055	<0.0055	<0.011	<0.028	<0.0055	<0.0055	<0.0055	<0.017
MP-22	7.0	44 YH	340 YH	200 YL	23 YL	<0.016	<0.016	<0.016	0.15	0.15	0.078	0.070	0.036	0.18
MP-23	6.5	<1.1	Z 6:1	<1.1	<5.7	<0.0057	<0.0057	<0.0057	<0.0114	<0.029	<0.0057	<0.0057	<0.0057	<0.017
	7.0	<1.1	<1.1	<1.1	<5.7	<0.0057	<0.0057	<0.0057	<0.0114	<0.029	<0.0057	<0.0057	<0.0057	<0.017

Notes:

<0.5]: Not detected at indicated concentration.

H: heavier hydrocarbons than indicated in standard

L : lighter hydrocarbons than indicated in standard

Y : sample exhibits fuel pattern which does not resemble standard

Z: sample exhibits unknown single peak or peaks

TPH-g: Total petroleum hydrocarbons as gasoline

TPH-JP5: Total petroleum hydrocarbons as jet fuel #5 TPH-d: Total petroleum hydrocarbons as diesel

TPH-mo: Total petroleum hydrocarbons as motor oil

from 6 of the 20 sampling locations at concentrations ranging from 0.6 micrograms per liter (μ g/L) (MP-19) to 71.9 μ g/L (MP-13). Table 4.2 summarizes groundwater analytical results from the March 1996 sampling. No mobile LNAPL was found.

BTEX was not detected in any of the three surface water samples collected. No sheens, odors, or other evidence of contamination was noted along the drainage channel or in the drainage ditch during sampling. However, 0.14 mg/L TPH-d was detected in the surface water sample collected from the small drainage ditch southwest of the site (sample location SW-02 on Figure 2.1). As discussed in Section 3.3.2.1, the surface water in the shallow drainage ditch is unlikely to be a discharge point for the shallow groundwater. Therefore, the low detection of TPH-d may be from surface runoff from the civil engineering support yard, which is located west of the site and upstream from this drainage ditch.

The TPH-g and BTEX concentrations measured in groundwater in March 1996 generally agree with previous sampling conducted in February 1995 (Appendix B). BTEX and TPH-g isopleths are shown on Figures 4.1 and 4.2. TMB concentrations are presented in Table 4.2. TMB results are presented because they are water-soluble fuel constituents that can be recalcitrant to biological degradation under anaerobic conditions; therefore, they can be used as tracer compounds in the calculation of anaerobic decay rates. Analytical results from the March 1996 investigation are discussed in the following subsections.

4.3.1.1 BTEX

The areal distribution of total dissolved BTEX in groundwater for the March 1996 TS is presented on Figure 4.1. As indicated by the 10-µg/L isopleth, the areal extent of the BTEX plume is estimated to be approximately 220 feet long and 180 feet wide and is elongated parallel to the direction of groundwater flow. The plume extends to the large drainage channel, which flows northeast to southwest, southeast of the former UST locations.

Because the plume has likely existed for a relatively long time period (more than 30 years), the plume has likely reached its maximum extent (i.e., the plume is no longer expanding). The maximum concentrations of BTEX were detected at MP-13 (71.1 μ g/L) and MP-14 (71.9 μ g/L), approximately 100 feet downgradient from the presumed source near the excavation pit. Although site access problems prevented installation of MPs to the southeast of the large drainage channel, the groundwater potentiometric surface map suggests that during the March 1996 sampling event the drainage channel was a discharge point for the shallow groundwater (see Section 3.3.2.1). Based on the March 1996 groundwater data, groundwater contaminants are not likely to underflow the channel; however, additional sampling east of the drainage channel and during the drier summer season would be required to confirm this conclusion.

Benzene was only detected at MP-14 at a concentration of $9.1\,\mu g/L$. This benzene concentration exceeds the California maximum contaminant level (MCL) for benzene of $1\,\mu g/L$ (California Code of Regulations, Title 22). MP-14 is downgradient of the source area; however, the relatively low BTEX concentrations and the short length of the plume prevents a determination of trends in benzene concentration in comparison to total BTEX with increasing distance from the source area.

ABLE 4.2

FUEL HYDROCARBON COMPOUNDS DETECTED IN GROUNDWATER/SURFACE WATER

UST Sites 10-494, 10-495, and 10-496 **MARCH 1996**

Intrinsic Remediation TS Beale AFB, California

Analyses		Tol Detroloum	Total Datroloum Undragarhan			Deare or By Campi ma	DTTT						
Allalyses.		Jiai relioieuli	II nyulocaloo	SI			BIEX			ŀ	her Volati	Other Volatile Aromatics	SS
Method:		USEPA S	USEPA SW8015M					USEPA SW	SW5030/SW8020A	020A			
							Ethyl-	Total	Total	1,2,3-	1,2,4-	1,3,5-	Total
Analyte:	IPH-g	1PH-JP5	p-Hd.I.	TPH-mo	Benzenc	Toluene	benzene	Xylenes	BTEX	TMB	TMB	TMB	TMB
Location	(mg/L)	u)	Ξ	(mg/L)	(hg/L)	(µg/L)	(µg/L)	(µg/L)	(μg/L)	(µg/L)	(μg/L)	(µg/L)	(µg/L)
MP-03	<0.050			<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MP-09	<0.050	0.71 YH	4.5 YH	3.5 Y	<0.5	5.0>	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
(dnp)	(<0.050)	(0.70 YH)	(4.5 YH)	(3.8 Y)	(<0.5)	(<0.5)	(<0.5)	(<0.5)	(<2.0)	(<0.5)	(<0.5)	(<0.5)	(<1.5)
MP-10	1.4	1.5 YH	4.3 YLH	2.2 YL	<0.5	<0.5	20	21.9	41.9	<0.5	3.6	7.8	11.4
MP-11	0.38 Y	0.13 Y	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
(dnp)	(0.41 Y)	(0.19 Y)	(0.094 YL)	(<0.30)	(<0.5)	(<0.5)	(<0.5)	(<0.5)	(<2.0)	(1.4)	(<0.5)	(<0.5)	(1.4)
MP-12	<0.005	n.a.	n.a.	n.a.	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MP-13	5.9	2.2 YH	3.5 YHL	1.6 YL	<2.5	9.2	22	39.9	71.1	120	<2.5	59	179
MP-14	4.0	1.9 YL	2.2 YL	0.31 YL	1.6	30	01	22.8	71.9	24	24	36	84
MP-15	0.13 Y	n.a.	n.a.	n.a.	<0.5	9.0	0.7	8.0	2.1	<0.5	<0.5	<0.5	<1.5
MP-17	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MP-18	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MP-19	<0.050	n.a.	n.a.	n.a.	<0.5	9.0	<0.5	<0.5	9.0	<0.5	<0.5	<0.5	<1.5
MP-20	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MP-21	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MP-22	0.77 Y	0.34 Y	0.32 Y	<0.30	<0.5	<0.5	<0.5	2.2	2.2	<0.5	<0.5	<0.5	<1.5
(dnb)	(0.79 Y)	(n.a.)	(n.a.)	(n.a.)	(<0.5)	(<0.5)	(1.8)	(<0.5)	(1.8)	(<0.5)	(<0.5)	(2.2)	(2.2)
MP-24	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MP-25	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MP-26	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MP-27	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MW-01	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
MW-02	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
SW-01	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
SW-02	<0.050	<0.050	0.140 Y	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
(dnp)	(<0.050)	(<0.050)	(0.120 Y)	(<0.30)	. (<0.5)	(<0.5)	(<0.5)	(<0.5)	(<2.0)	(<0.5)	(<0.5)	(<0.5)	(<1.5)
SW-03	<0.050	<0.050	<0.050	<0.30	<0.5	<0.5	<0.5	<0.5	<2.0	<0.5	<0.5	<0.5	<1.5
Notes:								4					

Notes:

<0.5]: Not detected above given concentration.

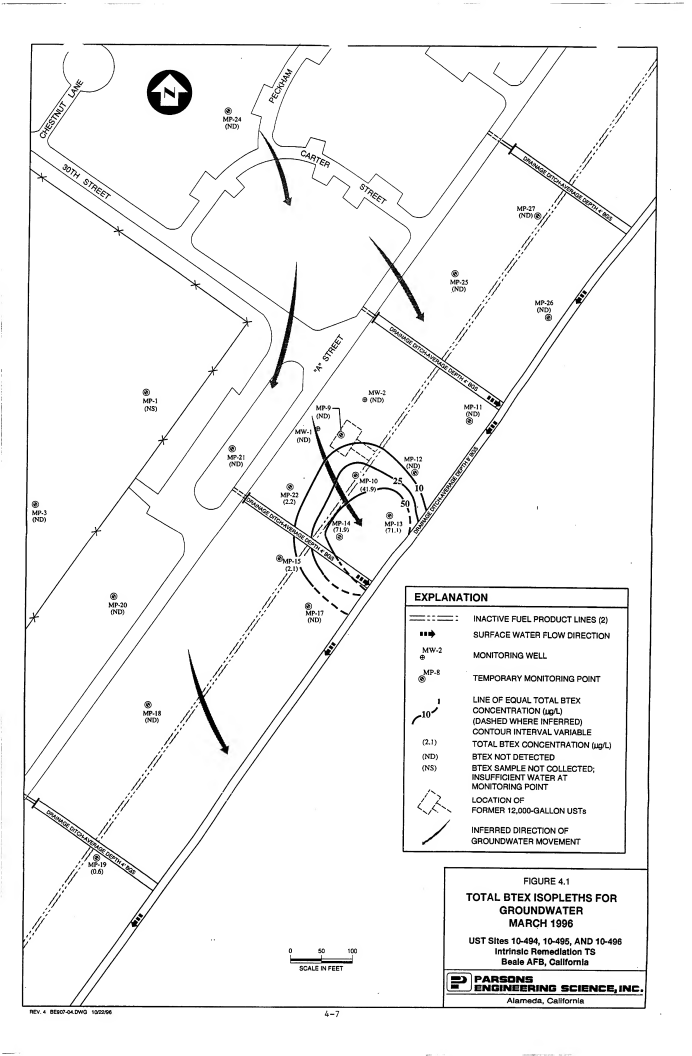
: Not analyzed. n.a.

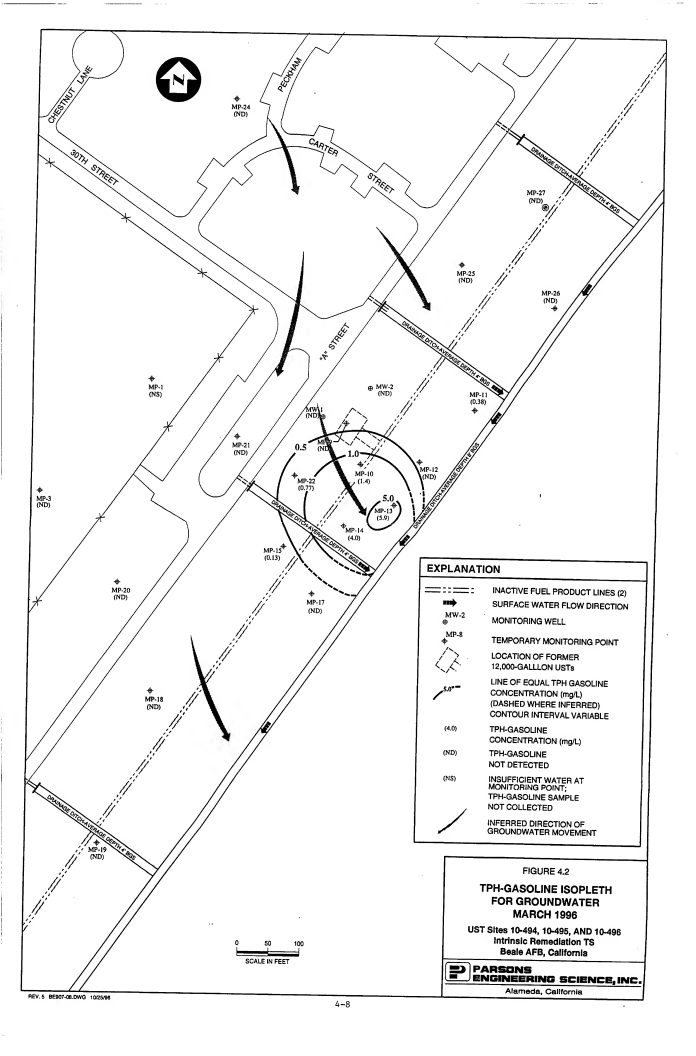
(dup): Primary analysis results followed in parentheses

[I]: heavier hydrocarbons than indicated in standard L: lighter hydrocarbons than indicated in standard

Y: sample exhibits fuel pattern which does not resemble standard

by duplicate analysis results.





Dissolved toluene, ethylbenzene, and total xylenes concentrations did not exceed their California MCLs of $150\,\mu\text{g/L}$, $750\,\mu\text{g/L}$, and $1,750\,\mu\text{g/L}$, respectively, at any sampled location. Total xylene concentrations exceeded the taste and odor threshold of $17\,\mu\text{g/L}$ at locations downgradient of the source area (MP-10, MP-13, and MP-14); toluene and ethylbenzene concentrations did not exceed their taste and odor thresholds at any location. Although toluene was detected slightly above the detection limit at MP-19 (0.6 $\mu\text{g/L}$), this detection appears to be unrelated to the plume at the UST Site based on data from the other MPs.

Because the plume extends to the large drainage channel, and as discussed in Section 3.3.2.1, this drainage channel is probably the point of groundwater discharge to surface water, the surface water historically may have received some mass of fuel-related contaminants from the site. An estimate of the BTEX mass released into the channel is made in Section 5.6 based on results of the calibrated groundwater model. The current lack of detectable concentrations of BTEX in surface water samples collected during this study indicates that the current rate of mass discharge into the surface water is low relative to the rate of natural attenuation of these chemicals in the channel and the volume of water flowing in the channel.

4.3.1.2 Total Petroleum Hydrocarbons

The areal distributions of TPH-g in groundwater for the TS sampling in March 1996 is presented on Figure 4.2. The shape of the TPH-g plume is nearly identical to that of the BTEX plume. Detected concentrations of TPH-g ranged from 0.13 mg/L at MP-15 to 5.9 mg/L at MP-13 (Table 4.2). TPH-g concentrations exceeded the taste and odor threshold at several locations downgradient and crossgradient of the source area (Table 4.2 and Figure 4.2). As with the soil analytical results, the laboratory reported some TPH quantitations as not matching the calibration standard, probably due to the highly weathered nature of the contamination. Detected TPH-g concentrations were generally greater than other TPH quantitations and less likely to be reported by the laboratory as not matching the calibration standard. These data are consistent with the historical evidence that gasoline was the fuel stored in (and released from) the USTs at the site.

4.3.1.3 Other Volatile Aromatics

The TMB isomers also were detected in MPs where BTEX and TPH-g were detected. TMBs are water-soluble fuel constituent and are used as tracers to estimate anaerobic decay rates for BTEX. The TMB results are discussed more fully in conjunction with the calculation of anaerobic decay rates (Section 5.3.5).

4.3.2 Inorganic Chemistry and Geochemical Indicators of BTEX Biodegradation

Microorganisms obtain energy for cell production and maintenance by facilitating thermodynamically advantageous redox reactions involving the transfer of electrons from electron donors to available electron acceptors. This results in the oxidation of the electron donor and the reduction of the electron acceptor. Electron donors at the UST Site are natural organic carbon and fuel hydrocarbon compounds. Fuel hydrocarbons are completely

degraded or detoxified if they are utilized as the primary electron donor for microbial metabolism (Bouwer, 1992). Electron acceptors are elements or compounds that occur in relatively oxidized states, and may include oxygen, nitrate, ferric iron, sulfate, and carbon dioxide.

The driving force of BTEX degradation is electron transfer, which is quantified by the Gibbs free energy of the reaction (ΔG°_{r}) (Stumm and Morgan, 1981; Bouwer, 1994; Godsey, 1994). The value of ΔG°_{r} represents the quantity of free energy consumed or yielded to the system during the reaction. Table 4.3 lists stoichiometry of the redox equations involving BTEX and the resulting ΔG°_{r} . Although thermodynamically favorable, most of the reactions involved in BTEX oxidation cannot proceed abiotically because of the lack of activation energy. Microorganisms are capable of providing the necessary activation energy; however, they will facilitate only those redox reactions that have a net yield of energy (i.e., $\Delta G^{\circ}_{r} < 0$).

Microorganisms preferentially utilize electron acceptors while metabolizing fuel hydrocarbons (Bouwer, 1992). DO is utilized first as the prime electron acceptor. After the DO is consumed, fuel-degrading anaerobic microorganisms typically use electron acceptors in the following order of preference: nitrate, ferric iron hydroxide, sulfate, and finally carbon dioxide.

Depending on the types and concentrations of electron acceptors present (e.g., nitrate. iron. sulfate, and carbon dioxide), pH conditions, and redox potential, anaerobic biodegradation can occur by denitrification, ferric iron reduction, sulfate reduction, or methanogenesis. Other, less common anaerobic degradation mechanisms such as manganese or nitrate reduction may dominate if the physical and chemical conditions in the subsurface favor use of these electron acceptors. Anaerobic destruction of the BTEX compounds is associated with the accumulation of fatty acids, production of methane, solubilization of iron, and reduction of nitrate and sulfate (Cozzarelli *et al.*, 1990; Wilson *et al.*, 1990). Environmental conditions and microbial competition ultimately determine which processes will dominate. Vroblesky and Chapelle (1994) show that the dominant terminal electron accepting process can vary both temporally and spatially in an aquifer with fuel hydrocarbon contamination.

Site groundwater data for electron acceptors strongly suggest that natural attenuation of hydrocarbons in the aquifer is occurring, and that aerobic biodegradation, denitrification, and iron reduction are the primary mechanisms through which remediation of the hydrocarbons in the aquifer is occurring. Geochemical parameters for site groundwater are discussed in the following sections.

4.3.2.1 Dissolved Oxygen

DO concentrations were measured at MWs and MPs during groundwater sampling in March 1996. Concentrations ranged from 0.15 to 8.40 mg/L. Table 4.4 lists the measured DO concentrations. Figure 4.3 is an isopleth map showing the distribution of DO concentrations in groundwater. Although comparison of Figures 4.1 and 4.3 shows graphically that areas with elevated total BTEX concentrations generally correlate with areas with depleted DO concentrations, the depleted DO concentrations may also be attributable to the former compost pile at the site. Therefore, DO may not have been an important electron acceptor at

TABLE 4.3

COUPLED OXIDATION REACTIONS FOR BTEX COMPOUNDS UST SITES 10-494, 10-495, AND 10-496 INTRINSIC REMEDIATION TS BEALE AIR FORCE BASE, CALIFORNIA

Coupled Benzene Oxidation Reactions	ΔG° _r (kcal/mole Benzene)	ΔG° _r (kJ/mole Benzene)	Stoichiometric Mass Ratio of Electron Acceptor to Compound
7.5 $O_2 + C_6 H_6 \Rightarrow 6 CO_{2,g} + 3 H_2 O$ Benzene oxidation /aerobic respiration	- 765.34	-3202	3.07:1
$6 NO_3 + 6 H^- + C_6 H_6 \Rightarrow 6 CO_{2,g} + 6 H_2 O + 3 N_{2,g}$ Benzene oxidation / denitrification	-775.75	-3245	4.77:1
$\frac{60 H^{+} + 30 Fe(OH)_{3,a} + C_{6}H_{6} \Rightarrow 6CO_{2} + 30 Fe^{2+} + 78H_{2}O}{Benzene \ oxidation / \ iron \ reduction}$	-560.10	-2343	21.5:1ª
$7.5H^{-} + 3.75SO_{4}^{2-} + C_{6}H_{6} \Rightarrow 6CO_{2,g} + 3.75H_{2}S'' + 3H_{2}O$ Benzene oxidation / sulfate reduction	-122.93	-514.3	4.61:1
$4.5 H_2O + C_6H_6 \Rightarrow 2.25 CO_{2,g} + 3.75 CH_4$ Benzene oxidation / methanogenesis	-32.40	-135.6	0.77:1

Coupled Toluene Oxidation Reactions	ΔG°r (kcal/mole Toluene)	ΔG°r (kJ/mole Toluene)	Stoichiometric Mass Ratio of Electron Acceptor to Compound
$9O_2 + C_6H_5CH_3 \Rightarrow 7CO_{2g} + 4H_2O$ Toluene oxidation /aerobic respiration	-913.76	-3823	3.13:1
$7.2NO_3 + 7.2H^- + C_0H_3CH_3 \Rightarrow 7CO_{2,g} + 7.6H_2O + 3.6N_{2,g}$ Toluene oxidation / denitrification	-926.31	-3875	4.85:1
$72 H^{-} + 36 \overline{Fe(OH)_{3,a}} + C_6 H_5 CH_3 \Rightarrow 7 CO_2 + 36 Fe^{2+} + 94 H_2 O$ $Toluene oxidation / iron reduction$	-667.21	-2792	21.86:1ª
$9H^{-} + 4.5SO_{4}^{2} + C_{6}H_{5}CH_{3} \Rightarrow 7CO_{2,g} + 4.5H_{2}S^{n} + 4H_{2}O$ Toluene oxidation / sulfate reduction	-142.86	-597.7	4.7:1
$5H_2O + C_6H_5CH_3 \Rightarrow 2.5CO_{2,\kappa} + 4.5CH_4$ Toluene oxidation / methanogenesis	-34.08	-142.6	0.78:1 ^b

TABLE 4.3 (continued)

COUPLED OXIDATION REACTIONS FOR BTEX COMPOUNDS UST SITES 10-494, 10-495, AND 10-496 INTRINSIC REMEDIATION TS BEALE AIR FORCE BASE, CALIFORNIA

Coupled Ethylbenzene Oxidation reactions	ΔG° _r (kcal/mole Ethyl-benzen e)	ΔG°r (kJ/mole Ethyl-benze ne)	Stoichiometric Mass Ratio of Electron Acceptor to Compound
$10.5O_2 + C_6H_5C_2H_5 \Rightarrow 8CO_{2,g} + 5H_2O$	-1066.13	-4461	3.17:1
Ethylbenzene oxidation /aerobic respiration			
$8.4 NO_3 + 8.4 H^- + C_6 H_5 C_2 H_5 \implies 8 CO_{2,g} + 9.2 H_2 O + 4.2 N_{2,g}$	-1080.76	-4522	4.92:1
Ethylbenzene oxidation / denitrification			
$84H^- + 42Fe(OH)_{3,a} + C_0H_5C_2H_5 \Rightarrow 8CO_2 + 42Fe^{2+} + 110H_2O$	-778.48	-3257	22:1ª
Ethylbenzene oxidation / iron reduction			
$10.5H^{-} + 5.25SO_{4}^{2} + C_{6}H_{5}C_{2}H_{5} \Rightarrow 8CO_{2,g} + 5.25H_{2}S'' + 5H_{2}O$	-166.75	-697.7	4.75:1
Ethylbenzene oxidation / sulfate reduction	20000		
$5.5H_2O + C_6H_3C_2H_3 \Rightarrow 2.75CO_{2g} + 5.25CH_4$	-39.83	-166.7	0.79:1 6
Ethylbenzene oxidation / methanogenesis			

Coupled m-Xylene Oxidation Reactions	ΔG°_{r} (kcal/mole m -xylene)	ΔG° _r (kJ/mole m-xylene)	Stoichiometric Mass Ratio of Electron Acceptor to Compound
$10.5O_2 + C_6H_4(CH_3)_2 \Rightarrow 8CO_{2,g} + 5H_2O$ m-Xylene oxidation /aerobic respiration	-1063.25	-4448	3.17:1
$8.4NO_3 + 8.4H^+ + C_0H_4(CH_3)_2 \Rightarrow 8CO_{2,g} + 9.2H_2O + 4.2N_{2,g}$ m-Xylene oxidation / denitrification	-1077.81	-4509	4.92:1
$84H^{-} + 42Fe(OH)_{3,u} + C_6H_4(CH_3)_2 \Rightarrow 8CO_2 + 42Fe^{2+} + 110H_2O$ $m-Xylene oxidation / iron reduction$	-775.61	-3245	22:1ª
$10.5 H^{-} + 5.25 SO_{4}^{2-} + C_{6}H_{4}(CH_{3})_{2} \Rightarrow 8CO_{2,g} + 5.25 H_{2}S'' + 5 H_{2}O$ $m-Xylene oxidation / sulfate reduction$	-163.87	-685.6	4.75:1
$5.5 H_2O + C_0H_4(CH_3)_2 \Rightarrow 2.75 CO_{2,g} + 5.25 CH_4$ m -Xylene oxidation / methanogenesis	-36.95	-154.6	0.79:1

a' Mass of ferrous iron produced during microbial respiration.

b/ Mass of methane produced during microbial respiration.

TABLE 4.4 GROUNDWATER AND SURFACE WATER **GEOCHEMICAL FIELD DATA**

MARCH 1996

UST Sites 10-494, 10-495, and 10-496 **Intrinsic Remediation TS** Beale AFB, California

	Dissolved	E_h		Ferrous		Carbon			
	Oxygen	(redox)	Total Iron	(Fe ⁺²) Iron	Mn	Dioxide	Temp.1	pН	EC
Location	(mg/L)	(mV)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(° C)	(units)	(µmhos/cm)
MP-03	3.40	394	0.06	ND	ND	58	15.0	6.40	235
MP-09			0.25	0.09	15.6	535			
(dup)	0.45	379	(0.35)	(ND)	(20)	(585)	18.0	6.65	4,150
MP-10	0.35	56	15.54	5.10	13.8	290	18.0	6.63	1,650
MP-11			0.09	0.07	2.9	42			
(dup)	0.20	245.7	(0.14)	(0.06)	(3.4)	(50)	15.0	6.76	200
MP-12	4.80	279.0	0.12	0.07	0.1	45	17.0	7.05	650
MP-13	0.40	128	7.86	6.84	26.4	210	16.0	6.98	1,100
MP-14	0.15	152.2	3.92	3.70	9.8	105	16.0	6.67	750
MP-15	2.60	302.3	0.25	0.16	1.8	46	n.a.	n.a.	n.a.
MP-17	8.00	426.5	0.06	ND	ND	30	16.0	6.85	180
MP-18	5.60	370.0	0.06	ND	ND	34	17.3	7.90	170
MP-19	n.a.	n.a.	0.12	ND	ND	20	n.a.	n.a.	n.a.
MP-20	4.20	422	0.12	0.07	0.2	60	16.5	6.42	235
MP-21	3.90	398.5	0.14	0.06	ND	50	19.0	7.51	215
MP-22			0.51	0.45	6.6	60			
(dup)	0.80	83.5	(0.46)	(0.34)	(8.7)	(55)	15.5	6.99	460
MP-24	8.40	392	0.13	ND	ND	28	16.0	6.34	145
MP-25	5.80	402.6	0.07	0.02	ND	34		6.65	145
MP-26	6.50		0.20	0.03	ND	36	15.5	6.85	
MP-27	6.10	372	0.10	ND	ND	33	15.2	6.59	137
MW-01	0.30	371	0.01	0.03	2.1	84	19.0	6.66	600
MW-02	1.80	.383	0.03	0.04	ND	26	19.9	8.32	220
SW-01	10.50	360	n.a.	n.a.	n.a.	n.a.	20.6	8.80	85
SW-02	11.50	354	n.a.	n.a.	n.a.	n.a.	24.2	8.18	280
SW-03	10.30	373	n.a.	n.a.	n.a.	n.a.	22.1	8.70	85

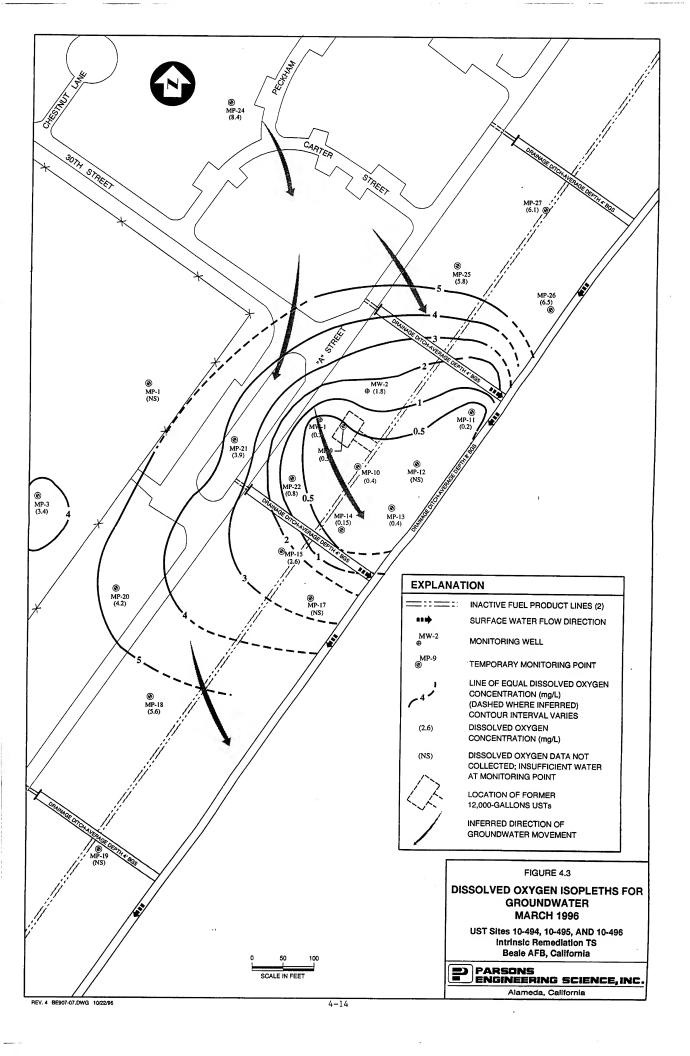
Notes:

ND: Not Detected. n.a.: Not analyzed.

MP-11 (dup): Primary sample analysis results followed by duplicate analysis results in parentheses.

¹ Measured with dissolved oxygen meter.

Eh: Oxidation - Reduction Potential relative to hydrogen.



the site in the past because the compost pile may have reduced available upgradient DO through elevated biological oxygen demand (BOD). However the removal of the compost pile in November 1995 and the average background DO concentration of 6.7 mg/L upgradient from the site (Figure 4.3) means that DO will be an important electron acceptor in the future.

MP-12 was not used in developing the DO isopleths. During purging, groundwater flow could not be sustained for a sufficient length of time to allow for DO to stabilize. A review of the boring log for MP-12 shows that the aquifer is very thin in this area of the site and is composed of silty, very fine to fine sand.

The stoichiometry of BTEX mineralization to carbon dioxide and water caused by aerobic microbial biodegradation is presented in Table 4.3. The average mass ratio of oxygen to total BTEX is approximately 3.14 to 1. This translates to the mineralization of approximately 0.32 mg of BTEX for every 1.0 mg of DO consumed. With a maximum observed background DO concentration of 6.7 mg/L, the shallow groundwater at this site has the capacity to assimilate 2.1 mg/L (2,100 μ g/L) of total BTEX through aerobic biodegradation. This may be a conservative estimate of the assimilative capacity of DO because microbial cell mass production was not taken into account by the stoichiometry presented in Table 4.3.

As a microbial population in the groundwater grows in response to the introduction of fuel hydrocarbons into the groundwater, some of the fuel is used not only to generate energy, but also to generate cell mass. When cell biomass production is accounted for, the mineralization of benzene to carbon dioxide and water is given by:

$$C_6H_6 + 2.5O_2 + HCO_3 + NH_4 \rightarrow C_5H_7O_2N + 2CO_2 + 2H_2O_3$$

This equation indicates that 5.0 fewer moles of DO are required to mineralize 1 mole of benzene when cell mass production is taken into account. On a mass basis, the ratio of DO to benzene is given by:

Benzene 6(12) + 6(1) = 78 gm

Oxygen 2.5(32) = 80 gm

Mass Ratio of Oxygen to Benzene = 80/78 = 1.03:1

On the basis of these stoichiometric relationships, 1.03 mg of oxygen are required to mineralize 1 mg of benzene, if cell mass production is taken into account. Similar calculations can be made for toluene, ethylbenzene, and the xylenes. On the basis of these calculations, approximately 0.97 mg of BTEX is mineralized to carbon dioxide and water for every 1.0 mg of DO consumed.

Although this process results in more efficient utilization of electron receptors, it is only applicable as the net cell mass of the microbial population continues to grow. Because groundwater contamination has been present at the UST Site for a number of years, biomass production within the center of the plume may have reached steady state. In such a case, the cell mass reaction equations would no longer apply. However, if the removal of the compost

pile (and its high BOD rates) results in a fresh influx of higher concentrations of DO into the area of the plume previously dominated by anaerobic processes, the higher rates of BTEX degradation could apply.

4.3.2.2 Nitrate/Nitrite

Concentrations of nitrate/nitrite [as nitrogen (N)] were measured in groundwater samples collected in March 1996. Table 4.5 summarizes measured nitrate and nitrite (as N) concentrations. Nitrate concentrations at the site ranged from 0.083 mg/L to 148 mg/L (as N). Nitrite was not detected in any sample. Figure 4.4 is an isopleth map showing the distribution of nitrate (as N) in groundwater. The maximum concentration of nitrate was detected at MP-9 (148 mg/L), which is located within the former excavation pit and compost pile. indicating that the compost pile was probably a significant source of nitrate in the groundwater. The nitrate concentration at MP-9 was two orders of magnitude higher than the average background concentration of 2.0 mg/L.

Comparison of Figure 4.4 with the BTEX plume on Figure 4.1 shows graphically that areas with elevated total BTEX concentrations have depleted nitrate concentrations. Although the former compost pile likely has been supplying nitrate to downgradient MPs, Figure 4.4 shows nitrate depletion in the BTEX plume to concentrations as low as 0.07 mg/L at MP-13. in the center of the BTEX plume. Concentrations of nitrate within the BTEX plume are below average background concentrations of approximately 2.0 mg/L, despite the nitrate source at the compost pile. These relationships provide strong evidence that anaerobic biodegradation of the BTEX compounds is occurring at the site through the microbially mediated process of denitrification.

In the absence of microbial cell production, the stoichiometry of BTEX mineralization to carbon dioxide. water, and nitrogen caused by denitrification is presented in Table 4.3. The average mass ratio of nitrate to total BTEX is approximately 4.9 to 1, or, when the nitrate is measured as N, the mass ratio of nitrate (as N) to total BTEX is 1.1 to 1. This translates to the mineralization of approximately 0.90 mg of BTEX for every 1.0 mg of nitrate (as N) consumed. With an observed average background nitrate concentration (excluding the concentration at MP-9) of 2.0 mg/L (as N), the shallow groundwater at this site has the capacity to assimilate 1.8 mg/L (1,800 μ g/L) of total BTEX during denitrification. This may be a conservative estimate of assimilative capacity for nitrate because microbial cell mass production has not been taken into account by the stoichiometry shown above (see Section 4.3.2.1) and because the former compost pile has not been incorporated into the calculation as a source of nitrate.

4.3.2.3 Ferrous Iron

Ferrous iron (Fe²⁺) concentrations were measured in groundwater samples collected in March 1996. Table 4.4 summarizes ferrous iron concentrations. Detected ferrous iron concentrations ranged from 0.02 mg/L to 6.84 mg/L. Figure 4.5 is an isopleth map showing the areal extent of ferrous iron in groundwater. Comparison of Figures 4.1 and 4.5 shows graphically that the area with elevated total BTEX concentrations also has elevated ferrous iron concentrations. Background concentrations of ferrous iron were generally below the

TABLE 4.5 GROUNDWATER AND SURFACE WATER GEOCHEMICAL ANALYTICAL DATA MARCH 1996

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

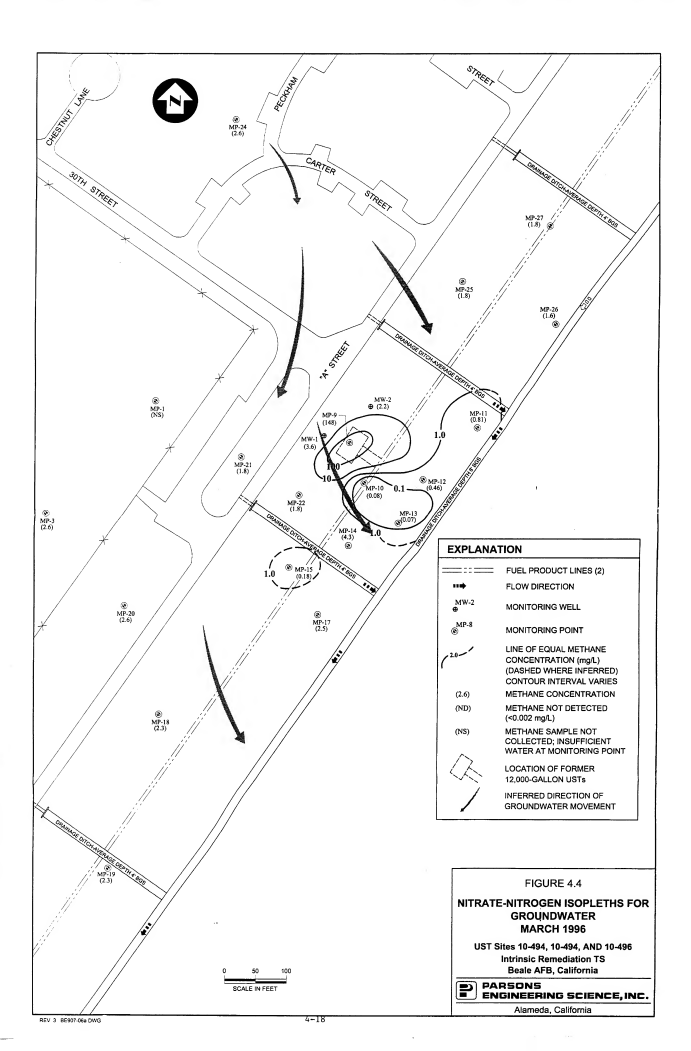
						Dissolved		
	Nitrate-	Nitrite-			Total	Ammonia-		
A 1			0.10.4	34.4		· ·	C1.1. 11.	TOC
Analyte:	Nitrogen	Nitrogen	Sulfate	Methane	Alkalinity	Nitrogen	Chloride	TOC
				RSKSOP-				
Method:	E300.0	E300.0	E300.0	175	E310.1	E350.3	E300.0	E415.1
Location	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)
MP-03	2.6	< 0.076	17.1	< 0.002	84.4	<0.8	25.7	1.4
MP-09	148	< 0.76	467	3.5	1,220	3.3	925	102
(dup)	(141)	(<0.76)	(458)	(3.9)	(1,250)	(1.9)	(885)	(103)
MP-10	0.083	<7.6	12.2	0.85	690	1.8	348	65.7
MP-11	0.81	< 0.076	7.0	0.176	104	<0.8	18	2.1
(dup)	(0.80)	(<0.076)	(7.0)	(0.167)	(103)	(<0.8)	(16.8)	(2.2)
MP-12	0.46	< 0.076	21.1	< 0.002	213	<0.8	136	5.2
MP-13	0.067	< 0.76	1.1	1.6	575	<0.8	195	36.8
MP-14	4.3	< 0.076	25.2	3.2	287	< 0.8	142	17.6
MP-15	0.18	< 0.076	11.4	0.023	n.a.	<0.8	84.4	6.6
MP-17	2.5	< 0.076	3.4	< 0.002	68.7	<0.8	20.3	1.5
MP-18	2.3	< 0.076	7	< 0.002	67.6	<0.8	10.9	2.2
MP-19	2.6	< 0.076	25.9	< 0.002	91.9	<0.8	103	3.2
MP-20	2.6	< 0.076	11.6	< 0.002	84.6	<0.8	25.1	1.2
MP-21	1.8	< 0.076	9.3	< 0.002	76.8	<0.8	23.3	1.8
MP-22	1.8	< 0.076	23.5	0.20	181	< 0.8	72.2	4.3
(dup)	(1.9)	(<0.076)	(23.5)	(0.22)	(186)	(<0.8)	(70.2)	(4.6)
MP-24	2.6	< 0.076	9.1	< 0.002	52.8	<0.8	11.2	<1
MP-25	1.8	< 0.076	5.8	< 0.002	60.4	<0.8	10.7	<1
MP-26	1.6	< 0.076	4.5	< 0.002	52.6	<0.8	8.1	<1
MP-27	1.8	< 0.076	5.2	< 0.002	59.4	< 0.8	10.2	<1
MW-01	3.6	< 0.076	26.8	< 0.002	177	<0.8	74.6	7.3
MW-02	2.2	< 0.076	8.4	< 0.002	86.4	<0.8	13.5	1.6
SW-01	0.15	< 0.076	1.6	n.a.	42.2	n.a.	2.3	n.a.
SW-02					92.2		.,	
(dup)	0.12	< 0.076	9.9	n.a.	(93.2)	n.a.	33	n.a.
SW-03	0.17	< 0.076	2.1	n.a.	40.3	n.a.	5.1	n.a.
Notes:								

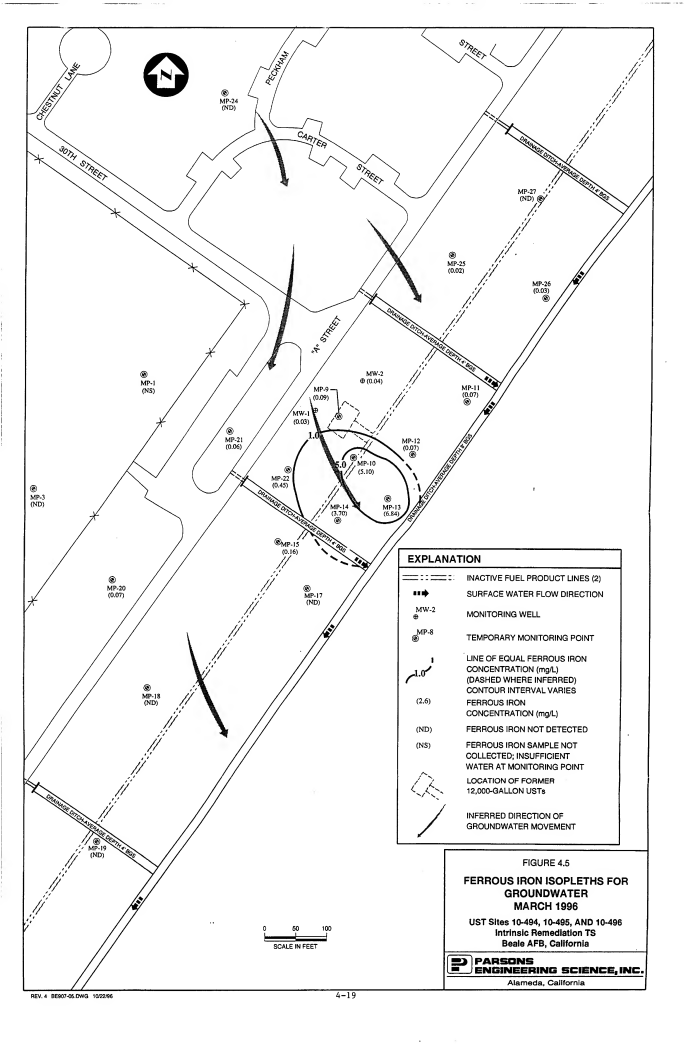
Notes:

<0.0002: Not detected above given detection limit.

n.a.: Not analyzed.

MP-11 (dup): Primary sample analysis results followed by duplicate analysis results in parentheses.





detection limit or at concentrations less than 0.10 mg/L. These relationships provide strong evidence that ferric iron hydroxide is being reduced to ferrous iron through anaerobic biodegradation of the BTEX compounds.

The stoichiometry of BTEX oxidation to carbon dioxide, ferrous iron, and water by iron reduction through anaerobic microbial biodegradation is presented in Table 4.3. On average, 37.5 moles of ferric iron hydroxide are required to metabolize one mole of total BTEX. Conversely, an average of 37.5 moles of ferrous iron are produced for each mole of total BTEX consumed. On a mass basis, this translates to approximately 21.8 mg ferrous iron produced for each 1 mg of total BTEX metabolized. Assuming a background ferrous iron concentration of zero and a maximum detected ferrous iron concentration of 6.84 mg/L at MP-13, the groundwater has the capacity to assimilate approximately 0.31 mg/L (310 μ g/L) of total BTEX through iron reduction. This is a conservative estimate of the assimilative capacity of iron because this calculation is based on observed ferrous iron concentrations. and not on the amount of ferric hydroxide available in the aquifer and solid soil matrix. Therefore, iron assimilative capacity could be higher.

Recent evidence suggests that the reduction of ferric iron to ferrous iron cannot proceed at all without microbial mediation (Lovley and Phillips, 1988; Lovley et al., 1991; Chapelle, 1993). None of the common organic compounds found in low-temperature, neutral, reducing groundwater could reduce ferric oxyhydroxides to ferrous iron under sterile laboratory conditions (Lovley et al., 1991). This means that the reduction of ferric iron requires microbial mediation by microorganisms with the appropriate enzymatic capabilities. Because the reduction of ferric iron cannot proceed without microbial intervention, the elevated concentrations of ferrous iron that were measured in the contaminated groundwater at the site are very strong indicators of microbial activity.

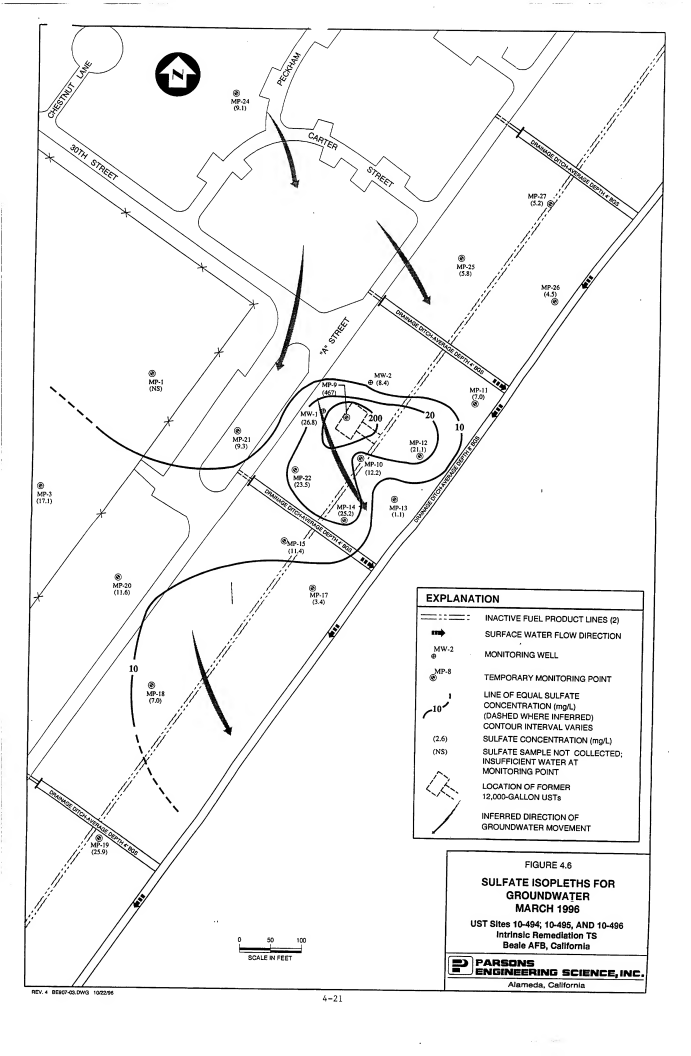
4.3.2.4 Sulfate

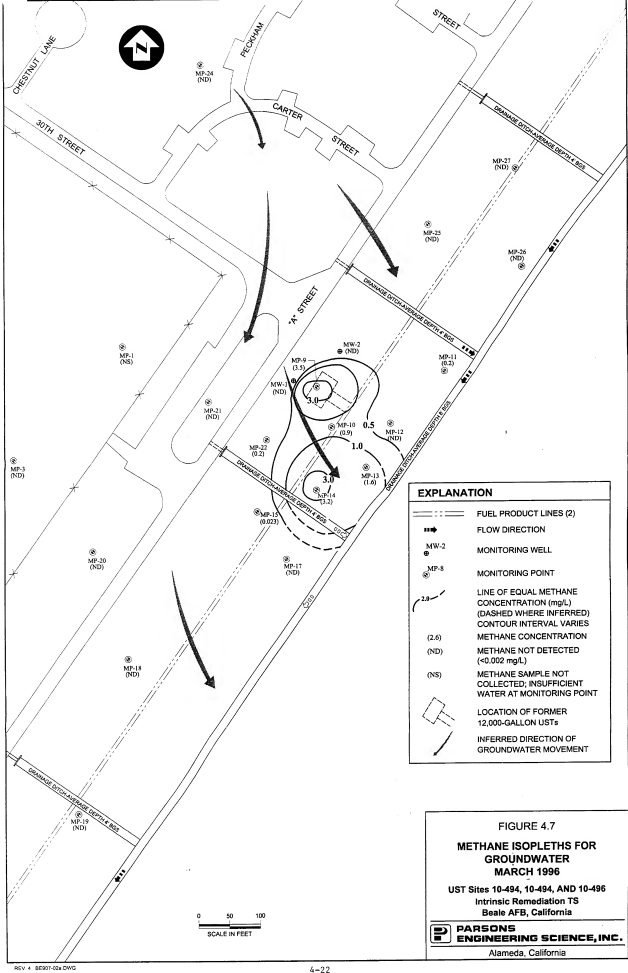
Sulfate concentrations in groundwater were measured at the site in March 1996. Table 4.5 summarizes sulfate concentrations. Sulfate concentrations ranged from 1.1 mg/L at MP-13 to 467 mg/L at MP-9. Figure 4.6 is an isopleth map showing the areal extent of sulfate in the groundwater.

The former compost pile appears to be a significant source of sulfate, similar to the results obtained for nitrate and discussed in Section 4.3.2.2. However, a comparison of sulfate concentrations relative to the BTEX plume does not show a definite pattern of sulfate depleted below background concentrations within the plume. Therefore, sulfate reduction is not considered an important process contributing to the natural attenuation of petroleum hydrocarbons at this site.

4.3.2.5 Methane

Methane concentrations were measured in groundwater samples collected in March 1996. Table 4.5 summarizes methane concentrations, which range from less than 0.002 to 3.5 mg/L. Figure 4.7 is an isopleth map showing the distribution of methane in groundwater. As





with nitrate and sulfate, the compost pile appears to be a significant source of methane to the groundwater.

However, elevated concentrations of methane also were detected at MP-14, where the highest concentration of BTEX was detected, and the methane concentration was depleted at MP-10. located along the groundwater flow path between the compost pile and MP-14. These results suggest that methanogenesis may be occurring at the site; however, the evidence is not conclusive because of the source of methane at the former compost pile. To be conservative, methanogenesis is not considered an important process contributing to the natural attenuation of petroleum hydrocarbons at this site.

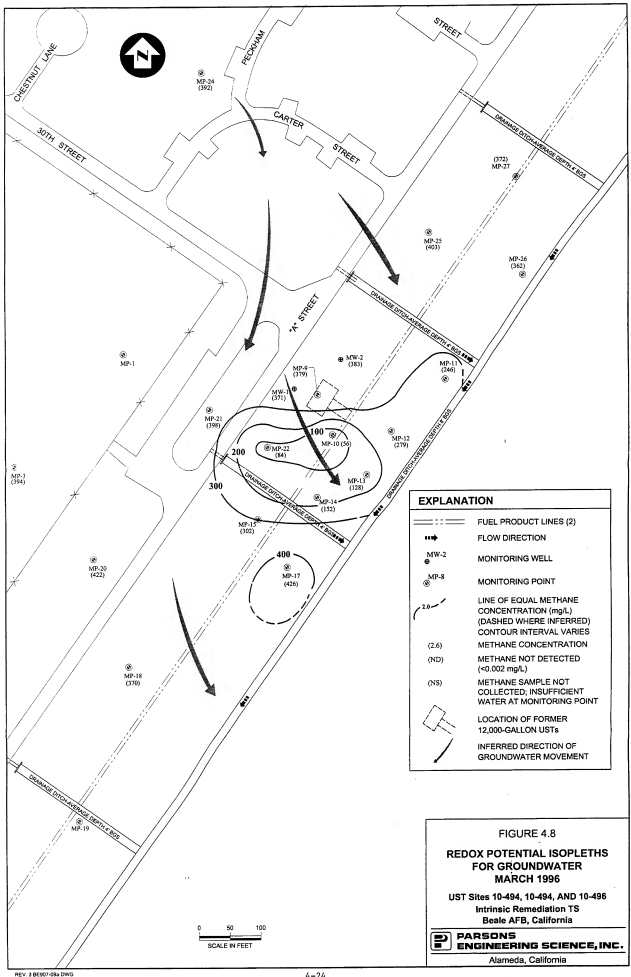
4.3.2.6 Reduction/Oxidation Potential

Redox potentials were measured in groundwater samples collected in March 1996. Redox potential is a measure of the relative tendency of a solution to accept or transfer electrons. The redox potential of a groundwater system depends on which electron acceptors are being reduced by microbes during BTEX oxidation. The redox potential at the site ranges from 56 millivolts (mV) to 426 mV, as measured relative to hydrogen. Table 4.4 summarizes available redox potential data. The areal extent of redox potentials is illustrated graphically on Figure 4.8. As expected, areas at the site with low redox potentials coincide with areas of high BTEX contamination, low DO and nitrate concentrations, and elevated ferrous iron concentrations.

The redox potentials measured at the site are higher than the theoretical optimum redox potentials for various electron acceptor reactions (Norris et al., 1994). This discrepancy is a common problem associated with measuring oxidizing potential using field instruments. Platinum electrode probes used in field instruments are not sensitive to the some of the redox couples (e.g., sulfate/sulfide). Many authors have noted that field measured redox data alone cannot be used to reliably predict the electron acceptors that may be operating at a site (Stumm and Morgan, 1981; Godsey, 1994; Lovley et al., 1994). Integrating redox measurements with analytical data on reduced and oxidized chemical species allows a more thorough and reasonable interpretation of which electron acceptors are being used to biodegrade site contaminants.

4.3.2.7 Alkalinity and Carbon Dioxide Evolution

Carbon dioxide is produced during the bioremediation of petroleum hydrocarbons. In aquifers that have carbonate materials as part of the matrix, carbon dioxide forms carbonic acid. which dissolves these minerals, increasing the alkalinity of the groundwater. An increase in alkalinity (measured as CaCO₃) above background conditions in an area with elevated BTEX concentrations can be used to infer the amount of petroleum hydrocarbon destroyed through aerobic respiration, denitrification, manganese reduction, ferric iron reduction, and sulfate reduction. In addition, carbon dioxide produced in these aerobic and anaerobic reactions can be cycled in the methanogenic reactions to continue BTEX biodegradation through methanogenesis.



Free carbon dioxide was measured in groundwater samples collected in March 1996 (Table 4.4). Carbon dioxide evolution above background concentrations is occurring as a result of biodegradation processes related either to the compost pile or BTEX; however, it is not possible to distinguish the two processes from each other. A direct estimate of the aquifer assimilative capacity based on carbon dioxide evolution is not possible because of the complex carbonate/bicarbonate balance.

Total alkalinity (as CaCO₃) was also measured in groundwater samples collected in March 1996 (Table 4.5). Alkalinity is a measure of the ability of groundwater to buffer changes in pH caused by the addition of biologically generated acids. Alkalinity also can be used in certain situations to estimate the assimilative capacity of groundwater (Wiedemeier et al., 1995). Total alkalinity measured at the site was relatively high and varied from 52.8 mg/L to 1,220 mg/L. Increases in alkalinity above background concentrations coincided with elevated carbon dioxide concentrations. Unfortunately, the production of carbon dioxide through elevated BOD from the compost pile prevented an estimate of the assimilative capacity of groundwater to degrade BTEX using alkalinity. However, the alkalinity concentrations and increases in carbon dioxide concentrations above background are supportive of an amenable environment for intrinsic biodegradation, and the alkalinity of site groundwater should be sufficient to buffer potential changes in pH caused by biologically mediated BTEX oxidation reactions.

4.3.2.8 pH

The pH was measured for groundwater samples collected in March 1996 (Table 4.4). The pH of a solution is the negative logarithm of the hydrogen ion concentration $[H^+]$. The optimal range of pH for BTEX-degrading microbes is between 6 and 8 standard units (Wiedemeier *et al.*, 1995). Only one groundwater sample, from MW-2, was slightly outside this range, at a pH of 8.32.

4.3.2.9 Temperature

The temperature of groundwater measured at the site in March 1996 is shown in Table 4.4. Temperature affects the type and growth rates of bacteria that can be supported in the groundwater environment. Temperatures in the shallow saturated zone varied from 15.0°C to 19.9°C. These are moderately warm temperatures for shallow groundwater, suggesting that bacterial growth rates should be stimulated.

4.3.3 Discussion

Three lines of evidence can be used to document the occurrence of natural attenuation: 1) geochemical evidence; 2) documented loss of contaminant mass at the field scale; and 3) laboratory microcosm studies. The first two lines of evidence (geochemical evidence and documented loss of contaminants) are used herein to support the occurrence of natural attenuation at the UST Site, as described in the previous sections. Because these two lines of evidence strongly suggest that natural attenuation is occurring at this site, laboratory microcosm studies were not deemed necessary.

Numerous laboratory and field studies have shown that hydrocarbon-degrading bacteria can participate in the degradation of many of the chemical components of jet fuel and gasoline. including the BTEX compounds (e.g., Jamison et al., 1975; Atlas, 1981, 1984, 1988; Gibson and Subramanian, 1984; Reinhard et al., 1984; Young, 1984; Bartha, 1986; Wilson et al., 1986, 1987, and 1990; Barker et al., 1987; Baedecker et al., 1988; Lee, 1988; Chiang et al., 1989; Grbic-Galic, 1989 and 1990; Cozzarelli et al., 1990; Leahy and Colewell. 1990; Altenschmidt and Fuchs, 1991; Alvarez and Vogel, 1991; Baedecker and Cozzarelli, 1991: Ball et al., 1991; Bauman, 1991; Borden, 1991; Brown et al., 1991; Edwards et al., 1991 and 1992: Evans et al., 1991a and 1991b; Haag et al., 1991; Hutchins and Wilson, 1991; Hutchins et al., 1991a and 1991b; Beller et al., 1992; Bouwer, 1992; Edwards and Grbic-Galic, 1992; Thierrin et al., 1992; Malone et al., 1993; Davis et al., 1994). Biodegradation of fuel hydrocarbons can occur when an indigenous population of hydrocarbon-degrading microorganisms is present in the aquifer and sufficient concentrations of electron acceptors and nutrients, including fuel hydrocarbons, are available to these organisms.

Comparison of BTEX, electron acceptor, and biodegradation byproduct isopleth maps (Figures 4.1 and 4.3 through 4.7) for the UST Site provides strong qualitative evidence of biodegradation of BTEX compounds. Isopleth maps suggest that the following electron acceptors are active in the biodegradation of BTEX compounds at the UST Site: DO, nitrate, and ferric iron (indicated by the presence of ferrous iron). Typically, zones of depleted oxygen, depleted nitrate, and elevated ferrous iron concentrations coincide through the core region of the BTEX plume because of the preferred order of electron receptor utilization as redox potential declines. At the UST Site, the sequence of electron acceptor use can be interpreted from the isopleths presented on Figures 4.3, 4.4, and 4.5 where the extent of the elevated or depleted concentrations generally decreases in order of decreasing energy yield.

4.3.4 Expressed Assimilative Capacity

The data presented in the preceding sections strongly suggest that mineralization of BTEX compounds is occurring through the microbially mediated processes of aerobic respiration, denitrification, and iron reduction. On the basis of the stoichiometry presented in Table 4.3, the expressed BTEX assimilative capacity of groundwater at the UST Site is approximately 4,200 μ g/L (Table 4.6). The primary mechanism is aerobic degradation, which accounts for approximately 2,100 μ g/L of expressed BTEX assimilative capacity. However, denitrification is an effective secondary mechanism, accounting for 1,800 μ g/L of expressed BTEX assimilative capacity. Sulfate reduction and methanogenesis may further contribute to the system's assimilative capacity; however, their impact is unclear as a result of the former composting operations at the site. Because of the removal of the compost pile, the groundwater geochemistry at the site can be expected to change within the next few years. It is recommended that the assimilative capacity of site groundwater be reevaluated during LTM at this site.

A closed system with 2 liters of water can be used to help visualize the physical meaning of assimilative capacity. Assume that the first liter contains no fuel hydrocarbons, but that it contains fuel degrading microorganisms and has an assimilative capacity of exactly "x" µg of fuel hydrocarbons. The second liter has no assimilative capacity; however, it contains fuel

TABLE 4.6 ASSIMILATIVE CAPACITY OF SITE GROUNDWATER

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

Electron Acceptor or Process	BTEX Assimilative Capacity (µg/L)
Dissolved Oxygen	2,100
Denitrification	1,800
Iron Reduction	310
Expressed Assimilative Capacity	4,210
Highest Observed Total BTEX Concentration	71.1

hydrocarbons. As long as these 2 liters of water are kept separate, biodegradation of the fuel hydrocarbons will not occur. If these 2 liters are combined in a closed system, biodegradation will commence and continue until the fuel hydrocarbons are depleted, the electron acceptors are depleted, or the environment becomes acutely toxic to the fuel-degrading microorganisms. Assuming a nonlethal environment, if fewer than "x" µg of fuel hydrocarbons were in the second liter, all of the fuel hydrocarbons will eventually degrade given a sufficient time; likewise, if greater than "x" µg of fuel hydrocarbons were in the second liter of water, only "x" µg of fuel hydrocarbons would ultimately degrade.

The groundwater beneath a site is an open system, which continually receives additional electron receptors from through the flow of the aquifer. This means that the assimilative capacity is not fixed as it is in a closed system, and therefore cannot be compared directly to contaminant concentration in the groundwater. Rather, the expressed assimilative capacity of groundwater is intended to serve as a qualitative tool. Although the expressed assimilative capacity at this site is greater than the highest measured total BTEX concentration, the fate of BTEX in groundwater and the potential impact to receptors is dependent on the relationship between the kinetics of biodegradation and the solute transport velocity (Chappelle, 1994). However, this significant expressed assimilative capacity is a strong indicator that an environment conducive to biodegradation of BTEX is present, and that the groundwater appears to have sufficient assimilative capacity to degrade the observed dissolved BTEX and limit plume migration.

SECTION 5

GROUNDWATER MODEL

5.1 GENERAL OVERVIEW AND MODEL DESCRIPTION

In order to help estimate degradation rates for dissolved BTEX compounds at the UST Site and to help predict the future migration of these compounds, Parsons ES modeled the fate and transport of the dissolved BTEX plume. The modeling effort had three primary objectives: 1) to predict the future extent and concentration of the dissolved contaminant plume by modeling the combined effects of advection, dispersion, sorption, and biodegradation; 2) to assess the potential for possible downgradient receptors to be exposed to contaminant concentrations greater than regulatory levels; and 3) to provide technical support for the evaluation of the RNA option at post-modeling regulatory negotiations. The model was developed using site-specific data and conservative assumptions about governing physical and chemical processes. Due to the conservative nature of the model input, the reduction in contaminant mass resulting from the process of natural attenuation is expected to exceed model predictions. This analysis is not intended to represent a baseline assessment of potential risks posed by site contamination.

The Bioplume II code was used to estimate the potential for dissolved BTEX migration and degradation by naturally occurring mechanisms operating at the UST Site. The Bioplume II model incorporates advection, dispersion, sorption, and biodegradation to simulate contaminant plume migration and degradation. The model is based upon the U.S. Geological Survey (USGS) Method of Characteristics (MOC) two-dimensional (2-D) solute transport model of Konikow and Bredehoeft (1978). The model was modified by researchers at Rice University to include a biodegradation component that is activated by a superimposed DO plume. On the basis of the work of Borden and Bedient (1986), the model assumes a reaction between DO and BTEX that is instantaneous relative to the advective groundwater velocity. Bioplume II solves the USGS 2-D solute transport equation twice, once for hydrocarbon concentrations in the aquifer and once for a DO plume. The two plumes are combined using superposition at every particle move to simulate the instantaneous, biologically mediated reaction between hydrocarbons and oxygen.

In recent years it has become apparent that anaerobic processes such as denitrification, iron reduction, sulfate reduction, and methanogenesis can be important BTEX degradation mechanisms (Grbic'-Galic', 1990; Beller et al., 1992; Edwards et al., 1992; Edwards and Grbic'-Galic', 1992; Grbic'-Galic' and Vogel, 1987; Lovley et al., 1989; Hutchins, 1991). Because geochemical evidence strongly supports the occurrence of anaerobic biodegradation processes at the UST Site (Section 4.3.2), the combined processes of aerobic and anaerobic biodegradation were considered in modeling BTEX fate and transport at the site. The

following subsections discuss in detail the input parameters, the model assumptions. the model calibration, and the simulation results.

5.2 CONCEPTUAL MODEL DESIGN AND ASSUMPTIONS

Prior to developing a groundwater model, it is important to determine if sufficient data are available to provide a reasonable estimate of aquifer conditions. In addition, it is important to ensure that any limiting assumptions can be justified. The most important assumption made when using the Bioplume II model is that oxygen-limited biodegradation is occurring at the site. The Bioplume II model assumes that the limiting factors for biodegradation are:

1) the presence of an indigenous hydrocarbon-degrading microbial population, and 2) sufficient background electron acceptor concentrations. Data and information presented in Sections 3 and 4 suggest that oxygen, nitrate, and ferric hydroxide are being used as electron acceptors for aerobic and anaerobic biodegradation.

On the basis of the data presented in Section 3, the aquifer is vertically defined by a layer of silty and/or gravelly fine- to coarse-grained sand. The majority of dissolved BTEX contamination preferentially migrates from the site in this sandy, alluvial layer that is 2 to 7 feet thick across most of the site. An underlying silty clay/clayey silt appears to restrict the vertical movement of groundwater, and the aquifer appears to be confined due to an overlying gravelly and sandy clay. The site is bounded on the east-southeast by a drainage channel that contains water year round and appears to intersect the upper coarse-grained layer. Groundwater enters the site from the northwest and flows across the site toward the drainage channel (Figure 3.10). The drainage channel appears to be a discharge point for the shallow groundwater as discussed in Section 3.3.2.1.

The aquifer was conceptualized and modeled as a shallow, confined aquifer composed of fine- to coarse-grained sand with some silt and clay fractions (Figures 3.8 and 3.9). The average saturated thickness of the aquifer was estimated at 4.5 feet. The model assumes horizontal flow with uniform velocity and direction. The use of a 2-D model is appropriate at the UST Site because the aquifer is relatively homogeneous and is bounded both above and below by clays and/or silts, which should inhibit vertical migration of dissolved BTEX contamination. Since a vertical mixing zone is not used in the model, the concentration front is assumed to be equivalent along the entire aquifer thickness.

BTEX was assumed to emanate from a fringe of soil contamination on the southern and northwestern sides of the former UST excavation based on analytical results from MP-10 (Table 4.1), soil vapor results for SVP-132 and SVP-142 (Appendix B and Figure 2.1), and previous observations of LNAPL during installation of SVP-132, SVP-142, and SB-6 (Appendix B). No additional fuel releases are expected at the site, and only BTEX contamination from the residual LNAPL contamination at the site was considered as a continuing source for the dissolution of BTEX into groundwater over time.

5.3 INITIAL MODEL SETUP

The setup for this model was based on available site data. Where site-specific data were not available (e.g., effective porosity), reasonable assumptions were made on the basis of widely accepted literature values for the types of materials comprising the shallow aquifer. The

following sections describe the basic model setup. Those Bioplume II model parameters that were varied during model calibration are discussed in Section 5.4.

5.3.1 Grid Design and Boundary Conditions

The maximum grid size for the Bioplume II model is limited to 20 columns by 30 rows. The dimension of each column and row can range from 0.1 to 999.9 feet. A 20- by 30-cell grid was used to model the UST Site. Each grid cell was defined as 20 feet wide by 20 feet long. The grid was oriented with the 30-cell dimension parallel to the large drainage channel in order to maximize the number of cells available for the model domain and minimize uneven boundary effects on cells near the drainage channel. The orientation results in a grid with a slightly offset direction from the dominant groundwater flow direction. However, this orientation is not expected result in a calibrated model with any additional sensitivity or error than that which would be imposed by reducing the number of cells or introducing uneven boundary effects. The grid encompasses the existing BTEX plume and covers an area of approximately 202,000 square feet, or 4.5 acres. The full extent of the model grid is indicated on Figure 5.1.

Model boundaries are mathematical statements that represent hydrogeologic boundaries, such as areas of specified head (i.e., surface water bodies or contour lines of constant hydraulic head) or specified flux. Hydrogeologic boundaries are represented by three mathematical statements that describe the hydraulic head at the model boundaries. These include:

• Specified-head boundaries (Dirichlet condition) for which the head is determined as a function of location and time only. Surface water bodies exhibit constant-head conditions. Specified-head boundaries are expressed mathematically as:

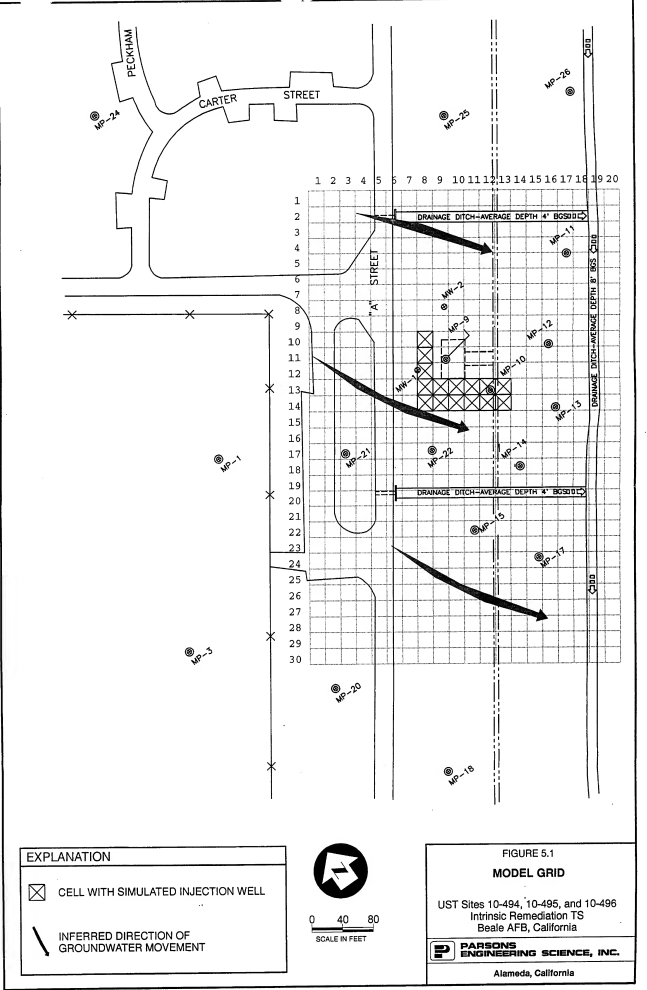
$$Head = f(x, y, z, t)$$

where f is the function symbol; x, y, and z are position coordinates; and t is time.

• Specified-flow boundaries (Neumann conditions) for which the mathematical description of the flux across the boundary is given. The flux is defined as a volumetric flow rate per unit area (i.e., cubic feet per square foot per day [ft³/ft²/day]). No-flow boundaries are a special type of specified-flow boundary and are set by specifying the flux to be zero. Examples of no-flow boundaries include groundwater divides and impermeable hydrostratigraphic units. Specified-flux boundaries are expressed mathematically as:

$$Flux = f(x, y, z, t)$$

 Head-dependent flow boundaries (Cauchy or mixed-boundary conditions) where the flux across the boundary is calculated from a given boundary head value. This type of flow boundary is sometimes referred to as a mixed-boundary condition because it is a combination of a specified-head boundary and a specified-flow boundary. Head-dependent flow boundaries are used to model leakage across semipermeable boundaries. Head-dependent flow boundaries are expressed mathematically as (Bear, 1979):



$$Flux = \frac{\left(H_0 - H\right)K'}{B'}$$

Where: H = Head in the zone being modeled (generally the zone containing the contaminant plume)

 H_0 = Head in external zone (separated from plume by semipermeable layer)

K' = Hydraulic conductivity of semipermeable layer

B' = Thickness of semipermeable layer

Natural hydraulic boundaries are modeled using a combination of the three types of model boundary conditions listed above. When possible, hydrologic boundaries such as surface water bodies, groundwater divides, contour lines, or hydrologic barriers should coincide with the perimeter of the model. In areas lacking obvious hydrologic boundaries, specified-head or specified-flux boundaries can be specified at the model perimeter if the boundaries are far enough removed from the contaminant plume that transport calculations are not affected. Bioplume II requires the entire model domain to be bounded by zero-flux cells (also known as no-flow cells), with other boundary conditions established within the subdomain specified by the no-flow cells.

Specified-head boundaries for the model were set in the northwestern corner of the model grid, where groundwater enters the site, and along the entire southeastern side of the model grid, coinciding with the surface water in the drainage channel. Typically in 2-D models, two rows of specified-head boundaries (along the top and bottom of the grid) are sufficient to simulate the flow of groundwater for sites that are not hydrogeologically complex or are bounded by adjacent lakes or streams. However, placement of specified-head cells on portions of three sides of the model grid for the UST Site was appropriate and necessary because of the surface water in the drainage channel and in order to correctly simulate the groundwater entering the site from the northwest. Using no flow boundaries for the remaining cells along the north, south, and west boundaries of the grid was appropriate because the potentiometric surface lines are approximately perpendicular to the model grid in these locations or are far enough removed from the area of interest (Figure 3.10).

The constant-head boundaries in the northwestern corner of the grid were chosen where the groundwater potentiometric surface map indicated constant-head contours (Figure 3.8). Constant-head values for the southeastern side of the grid were based on the surveyed surface water elevations (Table 3.2) and estimated upgradient and downgradient surface water levels in the drainage channel. The heads of the northwestern boundary were estimated to be from 129.75 to 129.5 feet above msl and represents the groundwater potentiometric surface in this portion of the site. The heads of the eastern boundary were estimated to vary from 126.09 to 124.5 feet above msl from north to south, respectively.

The base or lower boundary of the model is assumed to be no-flow and is defined by the underlying silt/clay at approximately 10 feet bgs. The upper model boundary is defined by the bottom of the overlying clay/silt aquitard located approximately 5 feet bgs.

5.3.2 Groundwater Elevation and Gradient

The March 1996 groundwater potentiometric surface map presented in Figure 3.10 was used to define the heads used as initial input into the Bioplume II model. The gradient range over the modeled area is approximately 0.0062 ft/ft to 0.018 ft/ft. Gradients are lowest upgradient from the contaminated source area and highest downgradient from the source area near MP-14 and the large drainage channel.

As discussed in Section 3.3.2.1, a slightly more southerly groundwater flow direction and lower hydraulic gradients are expected during the drier summer months. The groundwater potentiometric surface is also probably responsive to precipitation recharge, and will likely increase during the principal rainy season from November through April, and experience a subsequent decline during the main dry season from May through October. As described in Section 5.4.1, the model was calibrated to a reasonable match of the observed March 1996 potentiometric surface, which is expected to represent average site conditions.

5.3.3 BTEX Concentrations

As noted in Section 5.2, dissolved BTEX is assumed to enter groundwater at the UST Site from migration of recharge (precipitation) through soil containing residual LNAPL above the saturated zone (i.e., through leaching). The suspected areal extent of the residual LNAPL is estimated at approximately 6,000 square feet, surrounding the southern and northwestern portions of the former tank excavation pit (see Section 5.2). This estimate is based on observations of residual LNAPL and soil vapor sampling results from around the former excavation during previous investigations (Appendix B), and on the detection of TPH contamination at a concentration of 780 mg/kg at MP-10 (Table 4.2), located just outside the former excavation area. The area of residual LNAPL was accommodated in the model by 15 simulated injection wells, each within one 20-foot by 20-foot grid cell, to simulate the dissolution of BTEX compounds (Figure 5.1). The injection volume for the injection wells was set at a rate low enough that the hydraulic calibration for the model was not affected.

The total dissolved BTEX concentrations obtained from laboratory analytical results for each well and monitoring point location were used for model development. Table 4.1 presents dissolved BTEX concentration data, and isopleths of dissolved BTEX concentrations are shown on Figure 4.1. Section 5.3.4 describes the methods used to estimate the BTEX leaching rates from residual LNAPL used to calibrate the observed BTEX plume.

5.3.4 Source Estimate of Residual LNAPL Contamination

Groundwater can become contaminated when BTEX leaches from residual LNAPL contamination and into groundwater via infiltrating precipitation. As discussed in Section 3.4, recharge likely occurs on a seasonal basis even though the estimated annual ET rate is higher than the annual precipitation rate. An estimated 5 percent of annual rainfall (22.2 inches) is expected to reach the groundwater at the site. The remaining precipitation was assumed to be lost to runoff, evaporation, and ET. Evaporation was considered a large component of precipitation loss because the low-permeability clay/silt layer near the surface of the site (Figures 3.8 and 3.9) contribute to the pooling of water in depressions. The smaller drainage ditches north and south of the site carry some overland runoff into the larger

drainage channel; however, the site is relatively flat, as indicated by the surveyed elevations (Table 3.1). ET occurs across the site in the vegetated site areas, covered by open grassland. For these reasons, the estimated recharge rate of 5 percent per year was considered reasonable given site topography, stratigraphy, and ground cover.

The partitioning of total BTEX from soil into downward percolating precipitation is dominated by three conditions: 1) the concentrations of BTEX in the contaminated soil, 2) the concentrations of BTEX already present in the percolating precipitation, and 3) the contact time between the infiltrating water and the contaminated soil. It was assumed that rainwater percolating into the soil at the site is initially devoid of BTEX contamination. Contact time between infiltrating water and the soil contamination is dependent on the downward percolation rate. This parameter is difficult to estimate due to the influences of season, temperature, soil moisture, porosity, and soil type. To be conservative, it was assumed that all infiltrating water contacting areas of soil contamination were under equilibrium conditions. This means that infiltrating water would be in contact with contaminated soil for a sufficient length of time to allow maximum concentration of each of the BTEX compounds to dissolve and reach underlying groundwater.

On the basis of the work described by Bruce *et al.* (1991), the equilibrium saturation concentration of fuel hydrocarbon compounds in water can be estimated by:

$$C_w = C_f/K_{fw}$$

where:

 C_w = Concentration of a compound in aqueous solution

 C_f = Concentration of a compound in source soil

 K_{fw} = Partitioning coefficient

The fraction of BTEX in weathered residual LNAPL was estimated by computing the ratio of soil BTEX to soil TPH using the assumption that soil TPH concentrations were representative of the total fuel mass sorbed to the soil. This fraction was then compared to the fraction of BTEX in unweathered gasoline to estimate a concentration of BTEX in the weathered residual LNAPL in units of $\mu g/L$. The concentration was used to estimate the potential leaching rate of BTEX to groundwater in the areas of residual soil contamination at the site. The BTEX and TPH concentrations used for the leaching rate estimate were taken from sampling location MP-10 because it was adjacent to the former excavation pit and was the location where the maximum concentrations of BTEX and TPH were detected in soil. Therefore, the estimate is considered conservative for any remaining residual LNAPL at the site because the maximum soil concentration was used to compute the leaching rate for all the injection cells.

Based on the relatively small fraction of BTEX compounds remaining in the soil compared to that found in unweathered gasoline and the total concentration of TPH (Table 4.1), the residual LNAPL has been highly weathered. In order to simulate the higher rates of BTEX dissolution that occurred in the past, mass loading rates at the beginning of the model run were based on unweathered, fresh gasoline. The fraction of BTEX in the unweathered LNAPL was estimated from fresh gasoline fuels analysis data in the literature (Cline et al.,

1991; American Petroleum Institute, 1985); fresh gasoline has higher BTEX mass fractions than does the weathered LNAPL now present at the site. Mass loading rates at the end of the calibrated model run (present-day conditions) were based on the current highly weathered concentrations. The decreasing mass dissolution rates as the residual LNAPL weathered over time were estimated using a first-order decay constant. Use of a first-order constant was considered reasonable because the weathering of the residual LNAPL was assumed to consist qualitatively of both aerobic and anaerobic decay through vadose zone biodegradation as well as physical weathering and volatilization, and these processes were assumed to continue acting on the remaining fuel mass over time.

According to the partitioning coefficients that describe the relative solubility of each BTEX compound (see Appendix D), benzene had the fastest dissolution rate, followed by toluene, xylenes, and ethylbenzene, respectively. Appendix D contains the calculations for the residual LNAPL source term used for the groundwater model. The mass loading rate at the start of the model run (35 years before present) was estimated at approximately 1.23 milligrams per day per square foot (mg/day-ft²) of BTEX contamination reaching groundwater from residual LNAPL contamination. The mass loading rate at the end of the calibrated model run model (present-day conditions) was estimated at approximately 0.0048 mg/day-ft² of BTEX. Summed over a suspected area of residual contamination of 6,000 square feet, the residual BTEX source loading for the entire UST Site varied from 7,400 mg/day to 28 mg/day for initial conditions and present-day conditions, respectively. The first-order decay constant was calculated at 0.00044 per day.

As mentioned in Section 5.3.3, residual LNAPL mass loading rates were simulated via injection wells in 15 grid cells in the Bioplume II model (Figure 5.1). The BTEX source loading rates in the preceding paragraph translate to concentrations of 40.1 mg/L per cell and 0.16 mg/L per cell at an injected flow rate of 5 x 10⁻⁶ cubic foot per second (cfs), and these concentrations were used as inputs in the model to simulate LNAPL dissolution for initial conditions and present conditions, respectively. The simulated BTEX loading rates used during each year of the model run are discussed in Section 5.6.1.

5.3.5 Anaerobic Degradation Rates

Available data strongly suggest that anaerobic degradation is occurring at the site. Combined anaerobic processes account for approximately 50 percent of the assimilative capacity of site groundwater (Table 4.6). Anaerobic degradation must therefore be simulated with Bioplume II to make meaningful predictions. The Bioplume II model simulates anaerobic biodegradation by assuming that such degradation follows first-order kinetics. As with a large number of biological processes, anaerobic biodegradation can be estimated using a first-order rate constant and the equation:

$$\frac{C}{C_0} = e^{-kt}$$

Where:

C = Contaminant Concentration at Time t

 C_0 = Initial Contaminant Concentration

k = Coefficient of Anaerobic Decay (anaerobic rate constant)

t = time.

Two methods of calculating the anaerobic rate constants are currently available to quantify rates of biodegradation at the field scale and are applicable for use with available site data. The first method involves the use of a biologically recalcitrant compound found in the dissolved BTEX plume that can be used as a conservative tracer. The second method, proposed by Buscheck and Alcantar (1995), involves interpretation of a steady-state contaminant plume configuration and is based on the one-dimensional, steady-state analytical solution to the advection-dispersion equation presented by Bear (1979).

5.3.5.1 Trimethylbenzene/Tetramethylbenzene Tracer Method

In order to calculate anaerobic rate constants, the apparent degradation rate must be normalized for the effects of dilution caused by advective/dispersive processes and sorption. This can be accomplished by normalizing the concentration of each contaminant to the concentration of a component of gasoline (a tracer) that has similar sorptive properties but is fairly recalcitrant to biodegradation. Observed BTEX concentration data can be normalized to the TMB isomers 1,3,5-TMB, 1,2,4-TMB, and/or 1,2,3-TMB, tetramethylbenzene (TEMB), or to another tracer with similar physiochemical properties. The TMB and TEMB compounds can serve as good tracers because they can be recalcitrant under anaerobic conditions, and they have sorptive properties similar to the BTEX compounds (Cozzarelli *et al.*, 1990 and 1994). Thus, these compounds are assumed to respond similarly to the processes of advection, dispersion, and sorption without experiencing a reduction in concentration due to biodegradation.

The corrected concentration of a compound is the concentration of the compound that would be expected at one point (B) located downgradient from another point (A) after correcting for the effects of dispersion, dilution, and sorption between points A and B. One relationship that can be used to calculate the corrected contaminant concentration is:

$$C_{B,Corr} = C_B(TEMB_A/TEMB_B)$$

Where:

C_{B,Corr} = Corrected concentration of compound at Point B

 C_B = Measured concentration of compound at Point B

 $TEMB_A$ = Measured TEMB (or TMB) concentration at Point A TEMB_B = Measured TEMB (or TMB) concentration at Point B.

A log-linear plot of the corrected contaminant concentrations versus the travel time from the origin can be used to determine whether the data set can be described using a first-order exponential equation [i.e., the coefficient of determination (r^2) is greater than approximately 0.9]. When this occurs, the exponential slope can be used as the anaerobic rate constant.

Application of the tracer method to the UST Site was not possible because of the age of the residual LNAPL contamination and its highly weathered state. The declining mass loading

rate has caused source area contaminant concentrations to fall below those farther downgradient. In addition, the close proximity of the large drainage channel as a groundwater discharge point, and the small size of the plume, did not allow for collection of sufficient data points along the section of flowpath with decreasing contaminant concentrations for the application of the tracer method.

5.3.5.2 Method of Buscheck and Alcantar

Buscheck and Alcantar (1995) derive a relationship that allows calculation of first-order decay rate constants for steady-state plumes. This method involves coupling the regression of contaminant concentration (plotted on a logarithmic scale) versus distance downgradient (plotted on a linear scale) to an analytical solution for one-dimensional, steady-state. contaminant transport that includes advection, dispersion, sorption, and biodegradation. For a steady-state plume, the first-order decay rate is given by (Buscheck and Alcantar, 1995):

$$\lambda = \frac{v_c}{4\alpha_x} \left[\left[1 + 2\alpha_x \left(\frac{k}{v_x} \right) \right]^2 - 1 \right]$$

Where: λ = first-order decay rate

 v_c = retarded contaminant velocity in the x-direction

 α_x = dispersivity

 k/v_x = slope of line determined from a log-linear plot of contaminant concentration versus distance downgradient along flow path

The first-order decay rate includes biodegradation resulting from both aerobic and anaerobic processes; however, in the absence of oxygen, the first-order rate is equivalent to the anaerobic decay rate.

Application of the Buscheck and Alcantar method to the UST Site also was not possible for the same reasons discussed in Section 5.3.5.1. In addition, the plume must be at or close to a steady-state configuration for this method, which is unlikely to be the case at this site because of the limited BTEX source strength of the residual LNAPL and the recent compost pile influence on BOD and other geochemical parameters. Therefore, the anaerobic degradation rate also could not be estimated from field data using this method.

5.3.5.3 Selection of Anaerobic Decay Rate Constant

Because an anaerobic decay constant could not be determined from site field data, a review of recent literature was performed. Chapelle (1994) reported that at two different sites with anaerobic groundwater conditions, the anaerobic rate constants were both approximately 0.01 day⁻¹. Wilson *et al.* (1994) reported first-order anaerobic biodegradation rates of 0.05 to 1.3 week⁻¹ (0.007 to 0.185 day⁻¹); Buscheck *et al.* (1993) reported first-order attenuation rates in a range of 0.001 to 0.01 day⁻¹; and Stauffer *et al.* (1994) reported rate constants of 0.01 and 0.018 day⁻¹ for benzene and *p*-xylene, respectively.

An anaerobic rate constant of 0.00049 day⁻¹ was used as a starting value in the calibration of the Bioplume II model for this site. This rate is at the low end of values reported in the

literature and is the first-order decay constant used for an RNA study at a site at Travis AFB, California with similar geologic, hydrogeologic, and geochemical conditions (Parsons ES, 1996). This starting value was considered consistent with site conditions and also potentially conservative because a low anaerobic decay rate would result in less mass lost to anaerobic biodegradation.

5.4 MODEL CALIBRATION

Model calibration is an important component in the development of any numerical groundwater model. Calibration of the flow model demonstrates that the model is capable of matching hydraulic conditions observed at the site; calibration of a contaminant transport model superimposed upon the calibrated flow model helps verify that contaminant loading and transport conditions are being appropriately simulated. The numerical flow model presented herein was calibrated by altering transmissivity and constant-head boundary conditions in a trial-and-error fashion until simulated heads approximated observed field values within a prescribed accuracy. After calibration of the flow model, the numerical transport model was calibrated by estimating and adjusting solute loading and transport parameters in a trial-and-error fashion until the simulated BTEX plume approximated observed field values. Table 5.1 lists input parameters used for the modeling effort. Model input and output files are included in Appendix D.

5.4.1 Water Table Calibration

The groundwater at the UST Site was assumed to be influenced by continuous recharge and discharge at the constant-head cells within the model grid. The initial water levels at the constant-head cells and the transmissivity values were varied to calibrate the water table surface. The model was calibrated under steady-state flow conditions.

Hydraulic conductivity is an important aquifer characteristic that determines the ability of the water-bearing strata to transmit groundwater. Transmissivity is the product of the hydraulic conductivity and the thickness of the aquifer. An accurate estimate of hydraulic conductivity is important to help quantify advective groundwater flow velocities and to define the flushing potential of the aquifer and the quantity of electron-acceptor-charged groundwater that is entering the site from upgradient locations.

Saturated thickness data from previous reports, geologic logs, and water level measurements were used in conjunction with the hydraulic conductivity estimates from slug tests (see Sections 2.5 and 3.3.2.2) to create an initial transmissivity grid for the entire model. To better match heads in the model to observed values, the transmissivities were progressively varied in sections of the grid until the simulated water levels for cells corresponding to selected well locations matched the observed water levels as closely as possible. Figure 5.2 shows the calibrated water table.

The model was successfully calibrated using three hydraulic conductivities. In the area of the site where the slug tests were performed (MW-1) and for MPs which exhibited similar recharge rates, the average conductivity calculated from the slug tests (8.8×10^{-5} ft/s) was used. Upgradient from the site, the conductivity was doubled to 1.8×10^{-4} ft/s to reproduce the gentler gradient and potentiometric surface. A lower hydraulic conductivity of 1.0×10^{-5}

TABLE 5.1

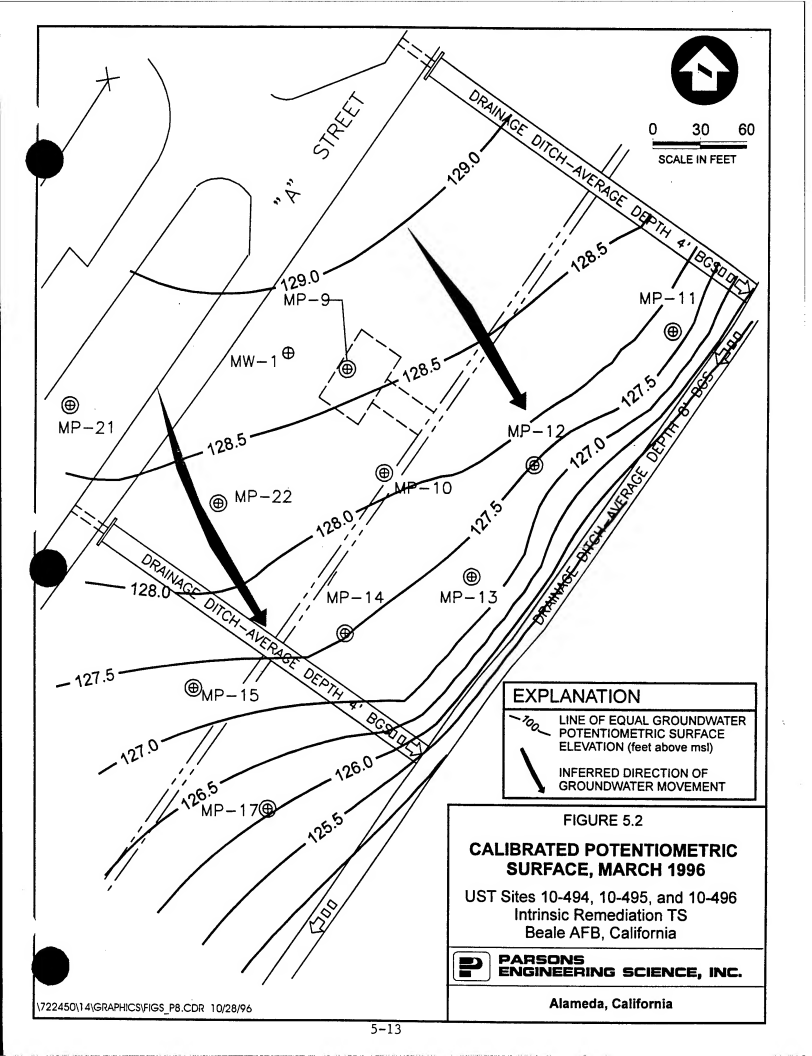
BIOPLUME II MODEL INPUT PARAMETERS

UST SITES 10-494, 10-495, AND 10-496 INTRINSIC REMEDIATION TS BEALE AFB, CALIFORNIA

Parameter	Description	Calibrated	Predictive Model	
	·	Model	(BEALESW)	
		(BEALECAL)		
NTIM	Maximum number of time steps in a pumping period	1	1	
NPMP	Number of Pumping Periods	36	50	
NX	Number of nodes in the X direction	20	20	
NY	Number of nodes in the Y direction	30	30	
NPMAX	Maximum number of Particles: NPMAX=	5245	5245	
	$(NX-2)(NY-2)(NPTPND) + (Ns^{a'})(NPTPND) + 250$			
NPNT	Time step interval for printing data	1	1	
NITP	Number of iteration parameters	7	7	
NUMOBS	Number of observation points	5	5	
ITMAX	Maximum allowable number of iterations in ADIP b	200	200	
NREC	Number of pumping or injection wells	15	15	
NPTPND	Initial number of particles per node	9	9	
NCODES	Number of node identification codes	2	2	
NPNTMV	Particle movement interval (IMOV)	0	0	
NPNTVL	Option for printing computed velocities	0	0	
NPNTD	Option to print computed dispersion equation coefficients	0	0	
NPDELC	Option to print computed changes in concentration	0	0	
NPNCHV	Option to punch velocity data	0	0	
NREACT	Option for biodegradation, retardation and decay	1	1	
PINT	Pumping period (years)	1	1	
TOL	Convergence criteria in ADIP	0.001	0.001	
POROS	Effective porosity	0.25	0.25	
BETA	Characteristic length (long. dispersivity; feet)	15	15	
S	Storage Coefficient	0	0	
		(Steady-State)	(Steady-State)	
TIMX	Time increment multiplier for transient flow	-	-	
TINIT	Size of initial time step (seconds)	-	-	
XDEL	Width of finite difference cell in the x direction (feet)	20	20	
YDEL	Width of finite difference cell in the y direction (feet)	20	20	
DLTRAT	Ratio of transverse to longitudinal dispersivity	0.1	0.1	
CELDIS	Maximum cell distance per particle move	0.5	0.5	
ANFCTR	Ratio of Tyy to Txx (1 = isotropic)	1	1	
DK	Distribution coefficient	0.32	0.32	
RHOB	Bulk density of the solid (grams/cubic centimeter)	1.65	1.65	
THALF	Half-life of the solute	-	-	
DEC1	Anaerobic decay coefficient (day ⁻¹)	0.00022	0.00022	
DEC2	Reaeration coefficient (day-1)	0	0	
F	Stoichiometric Ratio of Hydrocarbons to Oxygen	3.14	3.14	

a/ Ns = Number of nodes that represent fluid sources (wells or constant head cells)

b/ ADIP = Alternating-direction implicit procedure (subroutine for solving groundwater flow equation)



ft/s was used in areas around MP-12, MP-15, and MP-17 where recharge rates were very low during MP sampling activities. The average aquifer thickness of 4.5 feet determined from borehole logs (Appendix A) was used across the entire model grid. Water level elevation data from cells associated with the 11 groundwater monitoring locations within the model grid were used to compare measured and simulated heads for calibration. The root mean square (RMS) error is commonly used to express the average difference between simulated and measured heads. RMS error is the average of the squared differences between measured and simulated heads and can be expressed as:

RMS =
$$\left[\frac{1}{n}\sum_{i=1}^{n}(h_{m}-h_{s})_{i}^{2}\right]^{0.5}$$

Where:

n = the number of points where heads are being compared

 h_m = measured head value (feet above msl)

 h_s = simulated head value (feet above msl)

The RMS error between observed and calibrated values at the 11 comparison points was 0.243 foot, which corresponds to a calibration error of 9.1 percent. RMS error calculations are summarized in Appendix D. A plot of measured versus calibrated heads shows a random distribution of points around a straight line as shown in Appendix D. Deviation of points from a straight line should be randomly distributed in such a plot of results from computer simulations (Anderson and Woessner, 1992).

In solving the groundwater flow equation, Bioplume II establishes the water table surface and calculates an overall hydraulic balance that accounts for the numerical difference between flux into and out of the system. The hydraulic mass balance for the calibrated model was excellent, with greater than 99.99 percent of the water flux into and out of the system being accounted for numerically (a 0.0018 percent error). According to Anderson and Woessner (1992), a mass balance error of around 1 percent is acceptable, while Konikow (1978) indicates an error of less than 0.1 percent is ideal.

5.4.2 BTEX Plume Calibration

Model input parameters affecting the distribution and concentration of the simulated BTEX plume were modified so that model predictions matched dissolved total BTEX concentrations observed in March 1996. To do this, model runs were made using the calibrated steady-state hydraulic parameters coupled with the simulated introduction of contaminants at the injection wells.

Calibration of the fate and transport portion of a Bioplume II model generally requires that the contaminant distribution be known for at least two different times, one of which can be the initial release date. The time and volume of gasoline releases at the UST Site are not known; the USTs are simply known to have been removed prior to 1962.

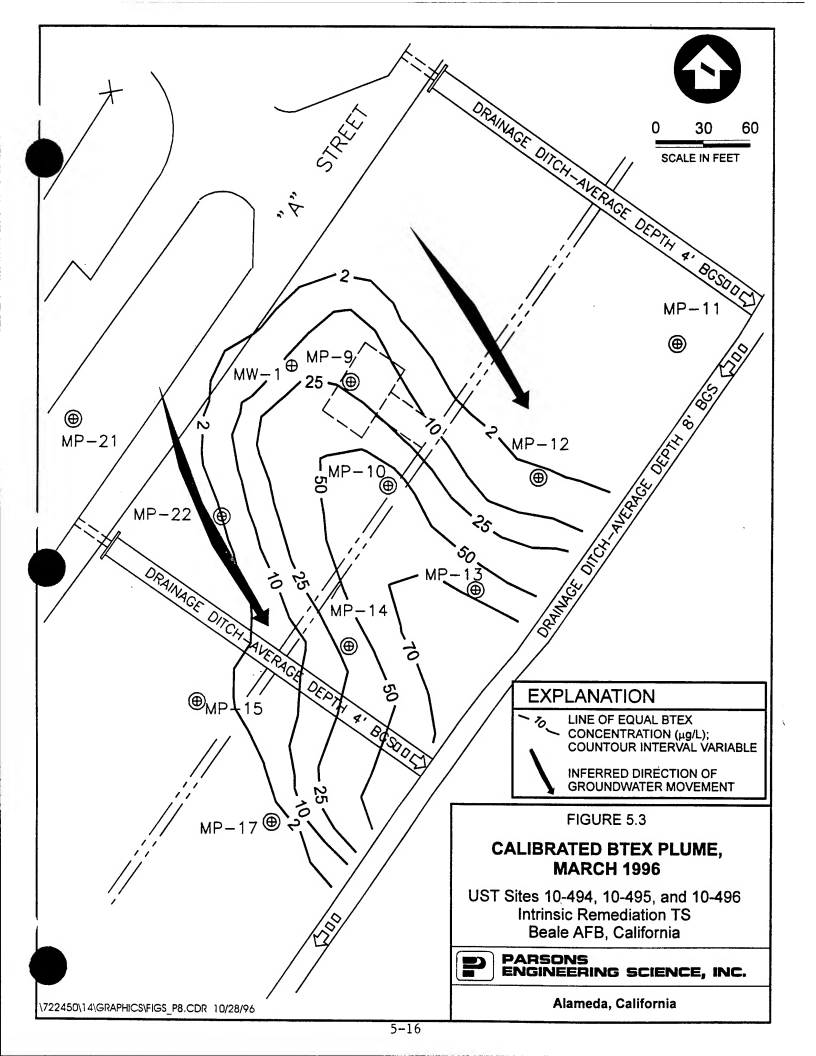
For the initial model run, it was assumed that the mass loading from the residual LNAPL in soil began in 1961. Mass loading rates were then decreased for the following 35 years of the model run (until 1996) using first-order decay to simulate weathering of the residual LNAPL source as discussed in Section 5.3.4. The estimated BTEX loading rates were applied to the 15 simulated injection wells of the model grid to reproduce the dissolution of BTEX from the residual LNAPL. While the term "injection well" suggests contaminants are being introduced at a point, Bioplume II assumes that contamination introduced at a well instantly equilibrates throughout the entire cell in which the well is located. The injection rate for the cells was set at 5 x 10⁻⁶ cfs, a value low enough that the flow calibration and water balance were not affected.

Although it is likely greater sources of BTEX contamination existed prior to UST removal activities (e.g., mobile LNAPL or higher amounts of residual LNAPL in the soil that was removed), model runs indicated that any concentration of dissolved BTEX that was input at the beginning of the model run would be either completely degraded or discharged to the drainage channel over the 35 year period of the model run. Because the extent of contamination at the time of the UST removal action is not known and because the results of the model were not affected by any pre-existing dissolved BTEX contamination, no initial BTEX concentrations were input into the final calibrated model. Therefore, mass estimates included in the model output files do not account for any initial mass which may have been in the aquifer prior to UST removal activities.

In addition to the injection of BTEX, Bioplume II allows the DO concentration to be specified at the beginning of the model simulation for the entire model grid. Bioplume II also allows constant DO concentrations in the water entering the model grid from constant-head cells. At this site, the presence of the compost pile within the former excavation pit posed a significant difficulty in modeling DO concentrations. The compost pile likely generated significant BOD from its initiation after UST removal activities and its removal in November 1995. However, there is no capability in Bioplume II to model an oxygen sink (i.e., the compost pile) separately from the contaminant (i.e., BTEX) being investigated; therefore, the calibrated model was run assuming that no aerobic biodegradation occurred between 1961 and 1996 (i.e., the initial DO concentration and the water entering the model grid were assumed to have DO concentrations of 0 mg/L).

By excluding aerobic biodegradation, the model is conservative because only anaerobic biodegradation (a slower process) is included. However, a significant disadvantage of this approach is that upgradient and crossgradient from the plume, modeled BTEX concentrations will be overestimated. DO is likely reaching cross-gradient locations and would be expected to limit the width of the plume. More importantly, DO is also likely reaching the locations of residual LNAPL upgradient from the compost pile at concentrations high enough to stimulate biodegradation of the leaching BTEX, especially in later years as the BTEX concentrations decrease.

Plume calibration was performed over a 35-year simulation period (1961 to 1996). Computed BTEX plume concentrations and configurations were compared to March 1996 BTEX data after 35 years of simulation time incorporating injection, transport, and degradation of contaminants in groundwater. The calibrated plume is shown in Figure 5.3.



The final calibrated model plume (year 35) was assumed to represent present-day (1996) conditions and compares favorably to the observed BTEX plume (Figures 4.1 and 5.3). As discussed above, the model overestimates the BTEX concentrations at upgradient (near MP-9 and MW-1) and crossgradient (near MP-12 and MP-17) locations.

The objective of the calibration was to achieve a modeled plume that equaled or exceeded the observed BTEX plume in terms of extent and concentration. The calibrated model successfully meets these objectives, as it reproduces both the observed areal extent of the plume and its contaminant concentrations. The calibrated BTEX plume accurately predicts a preferential flow of contamination toward MP-13 and MP-14 and the discharge into the large drainage channel.

The fact that the model concentrations in the source area and upgradient from the same area are slightly higher than observed concentrations means additional BTEX mass is accounted for in the model simulations and model predictions are conservative. Slight variations in shape between the model and the observed plume are likely due to subsurface heterogeneities in the hydraulic conductivity, reaeration coefficient, dispersivity, and retardation coefficient that are extremely difficult to identify in the field and to replicate in a discretized model domain.

5.4.2.1 Discussion of Parameters Varied During Plume Calibration

The transport parameters varied during plume calibration were dispersivity, the coefficient of retardation, and the coefficient of anaerobic decay. These parameters were generally varied with the intent of altering plume migration so that the observed plume extent was reproduced. While these parameters were varied with this common intent, each had a slightly different impact on the size and shape of the simulated plume.

5.4.2.1.1 Dispersivity

Much controversy surrounds the concepts of dispersion and dispersivity. Longitudinal dispersivity values for sediments similar to those found at the site range from 0.1 to 200 feet (Walton, 1988). Because longitudinal dispersivity is a function of scale, typically it is estimated based on the length of the contaminant plume. However, at this site the presence of the drainage channel limits the actual plume length. Therefore, the scale of observation used to estimate an initial dispersivity value was taken as twice the plume length (to include its mirror image across the channel). The longitudinal dispersivity was initially estimated at 5 feet, using approximately 1 percent (0.01) of twice the plume length of 240 feet.

During plume calibration, longitudinal dispersivity was varied between 5 and 25 feet. These values are relatively low compared to possible values cited in the literature (Walton, 1988). The use of low dispersivity values is a conservative estimate for modeling because low dispersivities cause less BTEX to be lost to dilution. The calibrated model uses a dispersivity of 15 feet. The ratio of transverse dispersivity to longitudinal dispersivity was maintained at 0.1 to help reproduce the plume width observed at the site.

5.4.2.1.2 Coefficient of Retardation

Retardation of the BTEX compounds relative to the advective velocity of the groundwater occurs when BTEX molecules are sorbed to organic carbon in the aquifer matrix. TOC analyses often are influenced by the presence of soil contamination, which may cause high soil TOC concentrations without necessarily reflecting an increase in the sorptive potential of soil. Therefore, TOC measurements used for retardation estimates should be taken from contaminant-free soils. Furthermore, TOC analysis should be taken across the phreatic surface rather than in the vadose zone to best represent the sorptive potential of saturated soils in the aquifer. The coefficients of retardation for the BTEX compounds were calculated based on measured TOC concentrations in the soils in and near the saturated zone at the site (Section 4.2.4), an assumed bulk density of 1.65 grams per cubic centimeter (g/cm³) (Freeze and Cherry, 1979), and published values of the soil sorption coefficients (K_{oc}) for the BTEX compounds as listed by Wiedemeier *et al.* (1995). The results of these calculations are summarized in Table 5.2.

The lower the assumed coefficient of retardation, the farther the BTEX plume will migrate downgradient. A higher assumed coefficient of retardation leads to more mass being retained nearer the source. Because the residual LNAPL is highly weathered at the site and the majority of the BTEX components found were ethylbenzene and xylenes (Table 4.2), which have higher retardation coefficients than benzene and toluene, an average retardation coefficient for all BTEX components (R=3.14) was used initially during plume calibration. This initial value also was used for the final calibrated model. An analysis of the sensitivity of the calibrated model to the coefficient of retardation also was performed and is discussed in Section 5.5.

5.4.2.1.3 Coefficient of Anaerobic Decay

The coefficient of anaerobic decay is a first-order rate constant used in Bioplume II to simulate the rate of use and replenishment of anaerobic electron acceptors in the groundwater. A coefficient of anaerobic decay of 0.00049 day⁻¹ was originally estimated, based on literature values and the first-order decay rate calculated from a site at Travis AFB, California, as discussed in Section 5.3.5.3. Use of an anaerobic decay coefficient is justified at this site because anaerobic decay mechanisms account for approximately 50 percent of the assimilative capacity at the site. The coefficient of anaerobic decay had a significant effect on the magnitude of the concentrations within the center of the plume and also controlled the amount of contaminant mass reaching the drainage channel.

Through iterative variation of the anaerobic decay coefficient and other select model parameters, the model was calibrated to 1996 analytical data. An anaerobic decay constant of 0.00022 day⁻¹ provided the best calibration and was used in the final calibrated model. This value was approximately one-half of the starting value and probably provided a better fit to site data because anaerobic decay processes at the Travis site, upon which the starting value was based, included sulfate reduction and methanogenesis, which are not considered important processes at the UST Site. An analysis of the sensitivity of the calibrated model to the anaerobic decay constant also was performed and is discussed in Section 5.5.

CALCULATION OF RETARDATION COEFFICIENTS

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

Retardation Coefficient, R	Porosity Maximum Minimum Average	1.05	1.11 2.44	1.28 4.54	1.24 4.06	1.25 4.19	1.21		1.17 3.14
	Maximum	2.15	3.76	7.80	88.9	7.13	6.18		5.11
Effective	Porosity	0.25	0.25	0.25	0.25	0.25	0.25		
Bulk Density 1	(kg/L)	1.65	1.65	1.65	1.65	1.65	1.65		
t K _d (L/kg)	Average	0.090	0.22	0.54	0.46	0.48	0.41		0.32
Distribution Coefficient K _d (L/kg) Bulk Density Effective	Minimum	0.007	0.017	0.042	0.036	0.038	0.032		0.025
	Maximum	0.17	0.42	1.0	0.89	0.93	0.79		0.62
Fraction Organic Carbon, f _{oc} ²	Average	0.001145	0.001145	0.001145	0.001145	0.001145	0.001145		
	Minimum Av	0.0022 0.00009 0.0	0.0022 0.00009 0.001145	0.0022 0.00009 0.0	0.00009 0.0	0.00000 0.0	0.00000		
	ш		0.0022		0.0022	0.0022	0.0022		
K _{oc} -	(L/kg)	79	190	468	405	422	357		
	Compound	Benzene	Toluene	Ethylbenzene	m-Xylene	o-Xylene	p-Xylene	BTEX	Compounds

Notes:

¹ From technical protocol document (Wiedemeier et al., 1995)

² From site data

 $K_d = f_{oc} \times K_{ow}$

5.5 SENSITIVITY ANALYSIS

The purpose of the sensitivity analysis is to determine the effect of varying model input parameters on model output. Based on the work of Rifai et al. (1988), the Bioplume II model is most sensitive to changes in the coefficient of aerobic decay (reaeration coefficient), the coefficient of anaerobic decay, and the hydraulic conductivity of the media and is less sensitive to changes in the retardation factor, porosity, and dispersivity. To evaluate the sensitivity of the calibrated model for the most sensitive parameters, the transmissivity (hence the hydraulic conductivity) and the coefficient of anaerobic decay were varied. In addition, a sensitivity analysis was performed for the retardation factor because of its effect on the amount of mass that potentially discharges into the drainage channel. Reaeration is assumed to be negligible at the site, and including it would only serve to make the model less conservative, so performing a sensitivity analysis was deemed unnecessary for this parameter.

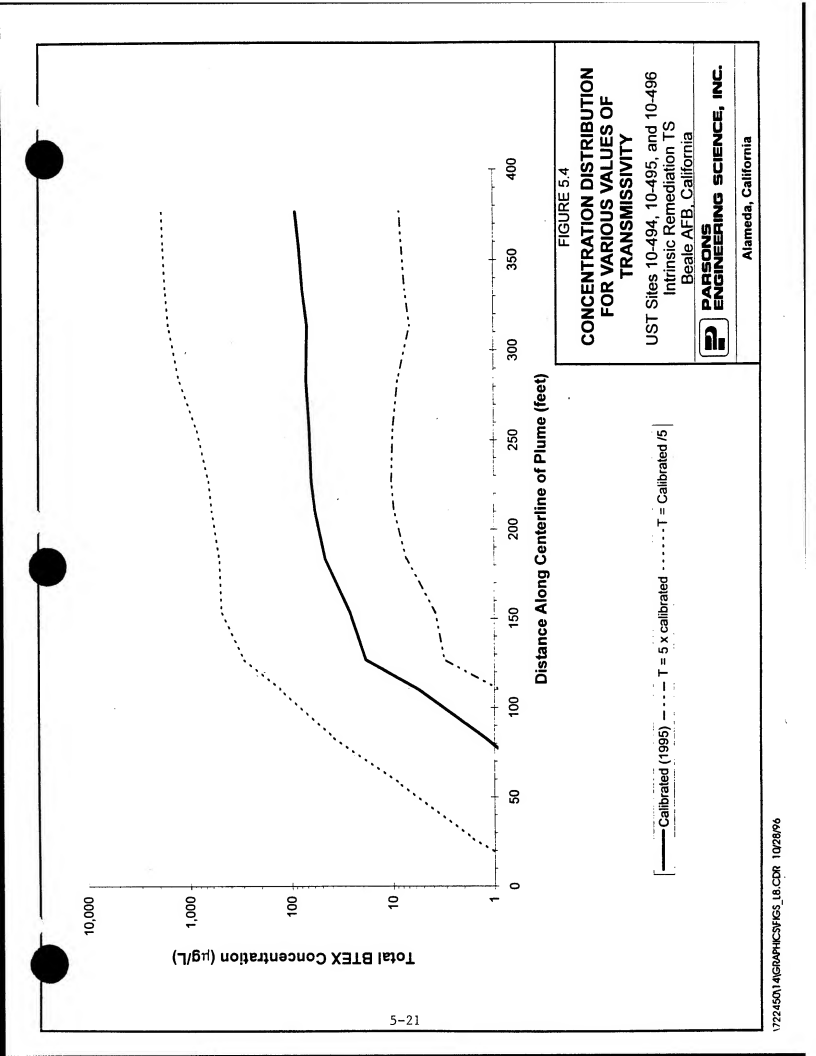
To perform the sensitivity analyses, the calibrated model was adjusted by systematically changing the aforementioned parameters individually and then comparing the new model runs to the original calibrated model. The sensitivity models were run for a 35-year period (the same duration used to achieve plume calibration in the original model) to assess the independent effect of each variable. As a result, six sensitivity runs of the calibrated model were made using the following variations:

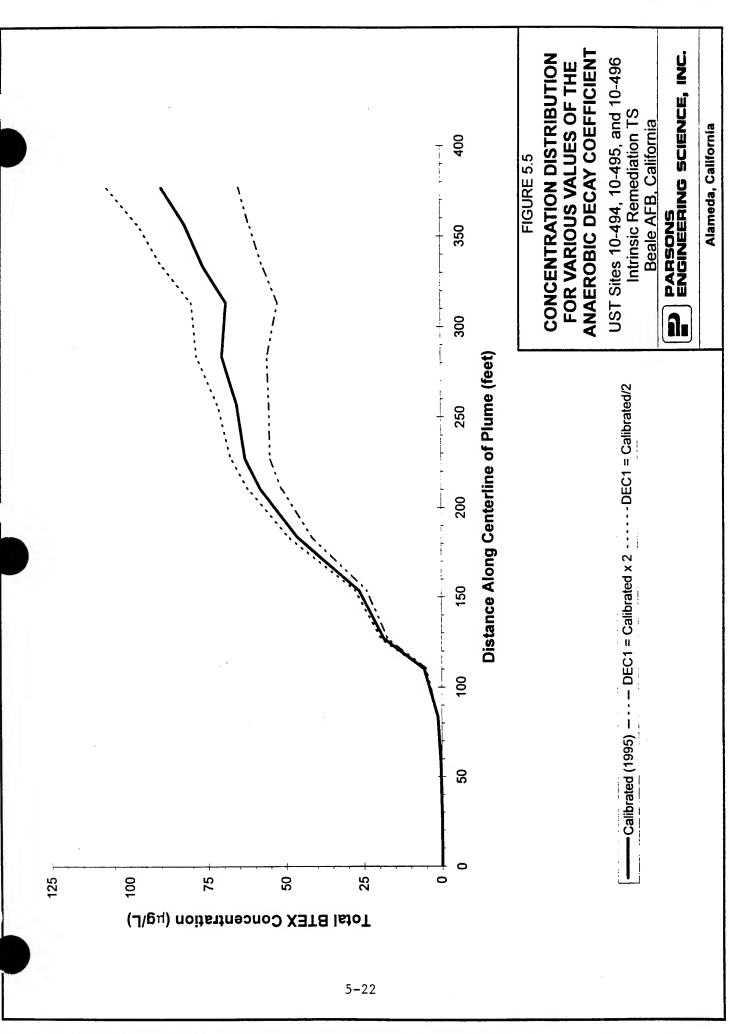
- 1. Transmissivity uniformly increased by a factor of 5;
- 2. Transmissivity uniformly decreased by a factor of 5;
- 3. Coefficient of anaerobic decay increased by a factor of 2;
- 4. Coefficient of anaerobic decay decreased by a factor of 2;
- 5. Coefficient of retardation increased by a factor of 2; and
- 6. Coefficient of retardation decreased by a factor of 2.

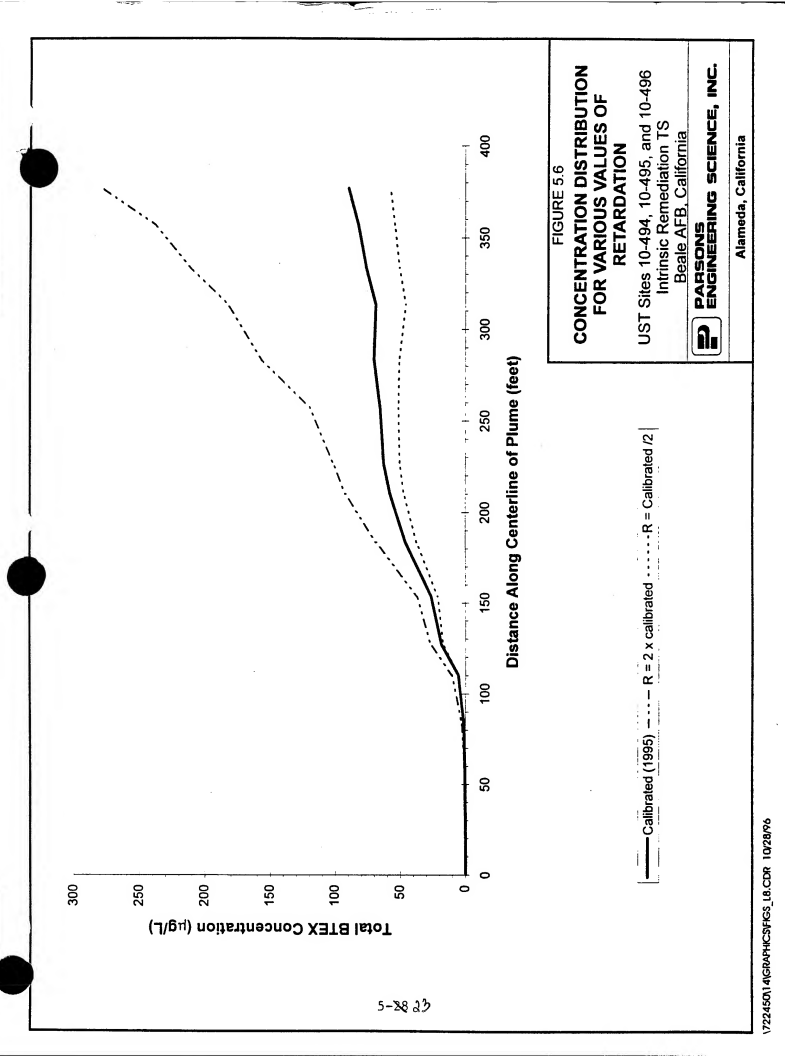
The results of the sensitivity analyses are shown graphically in Figures 5.4 through 5.6. These figures display the modeled BTEX concentrations versus distance along the centerline of the plume. This manner of displaying data is useful because BTEX concentrations are highest along the centerline and the plume is relatively symmetrical. Furthermore, the figures allow easy visualization of the changes in BTEX concentration caused by varying model input parameters.

The effects of varying transmissivity are shown on Figure 5.4. According to the work of Rifai et al. (1988), the Bioplume II model is particularly sensitive to variations in hydraulic conductivity. Higher values of hydraulic conductivity result in a faster-moving plume that may spread over a larger area and contains lower average BTEX concentrations. Lower values of hydraulic conductivity result in a slower-moving plume with a relatively small areal extent and higher average BTEX concentration.

Uniformly increasing the transmissivity by a factor of five significantly decreased the concentrations within the plume (by about one order of magnitude) due to the flushing effect







of faster groundwater velocities. The maximum concentration in the plume was only about $10~\mu g/L$ total BTEX, compared to the calibrated maximum of $89~\mu g/L$. Mass balance estimates made by the model indicated that because of the faster travel times to the drainage channel, approximately 92 percent of the total mass leaving the system (the mass leached from the residual LNAPL minus the mass still stored in the aquifer) discharged into the drainage channel. The remaining 8 percent of the mass was degraded anaerobically. In comparison, mass balance estimates for the calibrated model indicated that 67 percent of the total mass leaving the system discharged to the drainage channel and 33 percent was degraded anaerobically.

In contrast, uniformly decreasing the transmissivity by a factor of five significantly increased the concentrations within the plume (by between one and two orders of magnitude) due to the much longer residence times of the contaminants in the aquifer. Mass balance estimates made by the model indicated that because of the longer residence times and higher concentrations, only 12 percent of the total mass leaving the system discharged into the drainage channel. The remaining 88 percent of the mass was degraded anaerobically (higher concentrations lead to higher anaerobic mass loss because the decay is a first-order process). Therefore, the model indicates a somewhat counterintuitive result: higher concentrations within the aquifer lead to less mass discharging into the drainage channel over time when compared with the calibrated model. However, this conclusion is only a result of the site-specific conditions used for this model run (i.e., the selected anaerobic decay rate, the selected mass loading rate, and the distance to the drainage channel) and should not be construed to be a general conclusion. One can easily visualize situations where higher concentrations, lower anaerobic decay rates, and high transmissivities could lead to significant discharge to a surface water body.

These results show that, as expected, the model is very sensitive to transmissivity, suggesting that reasonable values were used in the calibrated model.

Figure 5.5 shows the effects of varying the anaerobic decay coefficient. Increasing this parameter by a factor of two results in more rapid degradation of dissolved BTEX and lower concentrations with increasing distance from the source. Although the concentrations are not significantly different with more rapid degradation, mass balance estimates made by the model indicate that over the 35-year period of the model run, only 45 percent of the total mass leaving the system discharged into the drainage channel (compared to 67 percent for the calibrated model). The remaining 55 percent of the mass was degraded anaerobically. The relative differences in concentration and total mass removed are not substantially higher because, during the initial periods of the model run, concentrations are so high and the drainage channel is so close to the source that substantial amounts of mass still discharge into the channel. At the end of the model run, the concentrations are so low that the amount of mass removed anaerobically is relatively small.

Conversely, decreasing the anaerobic decay coefficient by a factor of two also decreases the rate of degradation, resulting in an increase in the estimate of mass discharged to the channel (approximately 82 percent) and in the computed maximum BTEX concentration (to approximately $110 \, \mu g/L$). These results and the results discussed above show that the model

is only somewhat sensitive to the anaerobic decay coefficient with regard to concentration, and more sensitive to the anaerobic decay coefficient with regard to estimates of mass that discharge into the drainage channel. These results also indicate that reasonable values were used in the calibrated model.

The effects of varying the coefficient of retardation (R) are shown on Figure 5.6. Increasing R by a factor of 2 has a moderate effect on the contaminant concentration, while decreasing R by a factor of 2 only has a minor effect. An increase in sorptive capacity leads to a maximum BTEX concentration of 280 μ g/L (compared to 89 μ g/L for the calibrated model). As expected with higher concentrations and consistent with results discussed above for transmissivity changes, the mass of BTEX discharging to the drainage channel decreased from 67 percent for the calibrated model to 42 percent with the higher retardation coefficient.

The reason for the larger increase in concentration than decrease in concentration for the same change in the retardation coefficient is due primarily to the fact that the absolute value of the retardation factor cannot be less than one, and using a lower retardation factor simply means the contaminant velocity approaches the groundwater velocity. The retardation factor used for the calibrated model run was 3.14; decreasing it by a factor of 2 led to a retardation coefficient of 1.57, close to the limiting value of 1.

The results of the sensitivity analysis suggest that the calibrated model used for this report is reasonable. The calibrated model is most sensitive to transmissivity and slightly sensitive to the retardation coefficient and the anaerobic decay coefficient. Increasing the transmissivity greatly diminished the predicted BTEX concentrations, although the amount of mass discharged to the drainage channel also increased significantly. Lowering the values of transmissivity or increasing the retardation coefficient increased BTEX concentrations, but also increased the amount of mass anaerobically degraded and decreased the mass discharged into the drainage channel.

5.6 MODEL RESULTS

To predict fate and transport of dissolved BTEX compounds at the UST Site, the calibrated Bioplume II simulation (called BEALECAL) was run for a period of 50 years, 15 years beyond present-day conditions (Year 2011). This 50-year model run, called BEALESW (Beale Source Weathering), assumed the weathering of the residual LNAPL continued at the same first-order decay rate discussed in Section 5.3.4 and the degradation of dissolved BTEX contamination continued using the same anaerobic decay rate discussed in Section 5.4.2.1.3. The simulation assumed residual LNAPL will continue to exist at the site and no new removal actions or additional sources of BTEX will occur.

Table 5.3 shows the estimated reductions in BTEX loading rates applied to all years of the model run. These estimates are based on the calculations discussed in Section 5.3.4 and detailed in Appendix D. Input and output files for both the calibrated model and the source weathering model are provided on diskette in Appendix E.

Each of the injection periods assumes a BTEX injection rate lower than the previous period as a result of weathering using the first-order decay rate. To summarize, the weathering accounts for approximately 16 percent per year (%/yr) reductions in BTEX loading rates.

TABLE 5.3 ESTIMATED BTEX LOADING RATES FROM RESIDUAL LNAPL USED FOR MODEL RUNS

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

	Model	BTEX Loa	ading Rate
Year	Period	(mg/day-ft ²)	(ug/L-cell)
1961	1	1.2	40,100
1962	2	1.0	34,070
1963	3	0.88	28,940
1964	4	0.75	24,590
1965	5	0.64	20,890
1966	6	0.54	17,750
1967	7	0.46	15,080
1968	8	0.39	12,810
1969	9	0.33	10,880
1970	10	0.28	9,240
1971	11	0.24	7,850
1972	12	0.20	6,670
1973	13	0.17	5,670
1974	14	0.15	4,810
1975	15	0.13	4,090
1976	16	0.11	3,470
1977	17	0.090	2,950
1978	18	0.077	2,510
1979	19	0.065	2,130
1980	20	0.055	1,810
1981	21	0.047	1,540
1982	22	0.040	1,310
1983	23	0.034	1,110
1984	24	0.029	943
1985	25	0.024	801

	Model	BTEX Loa	ding Rate
Year	Period	(mg/day-ft ²)	(ug/L-cell)
1986	26	0.021	680
1987	27	0.018	578
1988	28	0.015	491
1989	29	0.013	417
1990	30	0.011	354
1991	31	0.0092	301
1992	32	0.0078	256
1993	33	0.0066	217
1994	34	0.0057	185
1995	35	0.0048	157
1996	36	0.0041	133
1997	37	0.0035	113
1998	38	0.0029	96
1999	39	0.0025	82
2000	40	0.0021	69
2001	41	0.0018	59
2002	42	0.0015	50
2003	43	0.0013	43
2004	44	0.0011	36
2005	45	0.00095	31
2006	46	0.00079	26
2007	47	0.00067	22
2008	48	0.00058	19
2009	49	0.00049	16
2010	50	0.00043	14

Notes:

- 1. Loading includes dissolution from residual LNAPL only (see Appendix D for calculation).
- 2. Loading does not account for sources which may have existed prior to UST removal (see Section 5.4.2).

Because the mass loading was modeled as a first-order process, mass loading into the aquifer is significantly higher during the initial periods of the model run, with approximately 90 percent of total BTEX removed from the total available mass in the first 15 years.

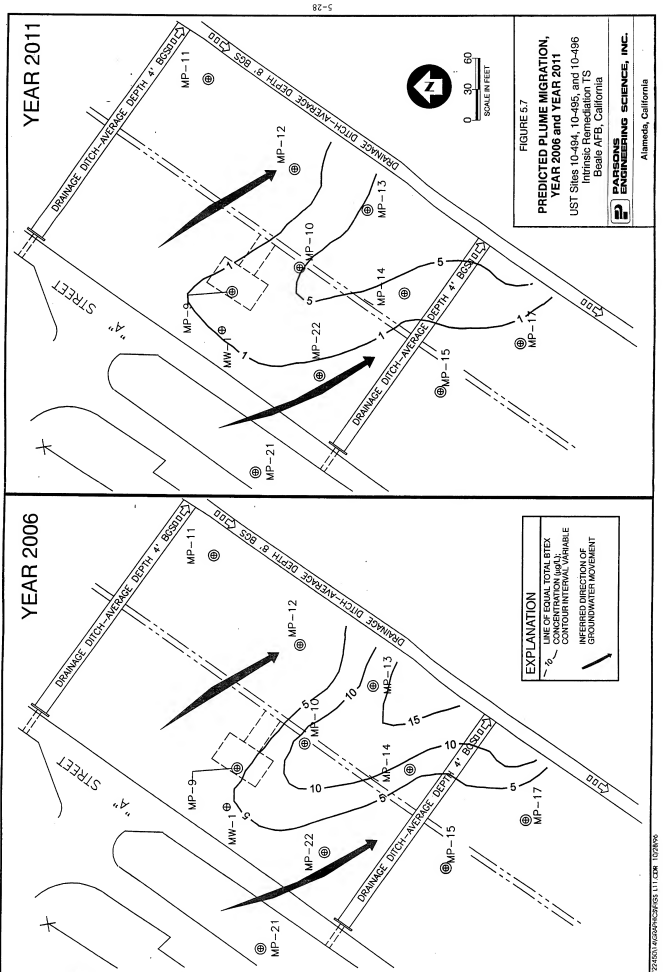
Using the conservative assumption that no aerobic decay occurs, over the 50-year period of the model run, approximately 67 percent of the mass leaving the system discharged to the drainage channel and 33 percent was degraded anaerobically. (Over the next 15 years, the model predicts that approximately 81 percent of the mass leaving the system will be discharged to the drainage channel and 19 percent will be degraded anaerobically.)

Based on the mass estimates made by the model, the amount of mass discharged to the drainage channel during 1995 was approximately 74 grams of BTEX. Using the estimated volumetric flow rate of water through the drainage channel of 2.5 cfs measured in March 1996. the estimated concentration of BTEX in the surface water would be less than $0.03~\mu g/L$, well below the analytical detection limit for BTEX compounds. This rough calculation provides only an order of magnitude estimate but indicates that the amount of mass discharging into the channel at the current time is probably not significant enough to adversely impact surface water. This conclusion also is consistent with the analytical results for surface water samples, discussed in Section 4.3.1.1. Although the flow in the drainage channel would be lower in the summer months leading to possibly higher surface water concentrations, the hydraulic gradient at the site is also expected to be lower in the summer leading to a lower discharge rate, and the two effects could serve to counteract each other.

Figure 5.7 shows the modeled plume for 2006 and 2011, 45 and 50 years from the start of the model run, respectively, and 10 and 15 years from present-day, respectively. After 10 years, modeled BTEX contours decrease to approximately 15 percent of their current concentrations. Based on the ratio of benzene to total BTEX found at MP-14, in 2006 benzene would be only slightly above its California MCL of 1.0 μ g/L. After 15 years, all areas of the plume are at or below 5 μ g/L total BTEX. After 15 years, both TPH and BTEX constituents also would be expected to be below their taste and odor thresholds.

Because the model continues to assume that no DO will be present in the aquifer, it represents a worst-case scenario and not necessarily the expected future site conditions. The removal of the compost pile in November of 1995 may allow some concentration of DO to flow into the site and rapidly reduce current and future BTEX mass. Because all BTEX concentrations at the site are below 72 μ g/L, only 230 μ g/L (0.23 mg/L) of DO would be required to completely mineralize BTEX based on the stoichiometric relationships presented in Section 4.3.2.1. With an upgradient DO concentration potentially as high as 8.4 mg/L, even a small contribution from upgradient groundwater would be sufficient to biodegrade all remaining BTEX at the site.

These results suggest that, under even the worst-case scenario, the observed BTEX plume will continue to attenuate over the next 10 to 15 years, and all groundwater at the site should be below California MCLs within approximately 10 years and below taste and odor thresholds within approximately 15 years. Rates of mass discharge into the drainage channel do not appear to be high enough to significantly impact surface water. In addition, even if a relatively small concentration of DO is available at the site in the future due to the recent



removal of the compost pile, the plume would attenuate much more rapidly and California MCLs and taste and odor thresholds could be reached within only a few years or less.

5.7 CONCLUSIONS AND DISCUSSION

A single conservative model scenario was used to predict BTEX attenuation and migration rates at the UST Site. Model BEALESW assumed weathering of residual LNAPL in soil using a first-order decay rate and mass dissolution to groundwater through leaching and desorption. The first-order decay rate was calculated by assuming that residual LNAPL initially consisted of fresh gasoline that degraded over a 35-year period to the highest measured TPH and BTEX concentrations found at the site during the 1996 investigation. Only anaerobic processes were assumed to occur at the site due to the presence of a compost pile, which probably acted as a significant sink for DO in the aquifer in the past.

The model predicted that over the next 10 to 15 years, even without any influx of DO. California MCLs and taste and odor thresholds would be met at the site with no engineered remediation. In addition, the model predicts that the current and future rates of mass discharge into the drainage channel are not expected to be high enough to significantly impact surface water.

Several conservative assumptions are built into the model. The use of these conservative model assumptions suggests that natural attenuation of BTEX contamination at the site may exceed model predictions. These conservative model assumptions include:

- 1. Aerobic biodegradation was not included for future predictions. Although not including aerobic processes for model calibration was appropriate because of the compost pile, the compost pile has been removed and it can be expected that some DO will begin to enter the site over the next few years.
- 2. The anaerobic decay coefficient used in the calibrated model is lower than common literature values for BTEX.
- 3. The stoichiometry used to determine the ratio between electron receptors and total BTEX assumed that no microbial cell mass was produced during the reactions. As discussed in Section 4.3.2, this approach may be too conservative by a factor of three.
- 4. An average coefficient of retardation for all BTEX compounds was used in the model simulations (R=3.1). The majority of the BTEX contaminant mass at the site is composed of toluene, ethylbenzene, and xylenes (Table 4.2). Retardation coefficients for toluene, ethylbenzene, and xylenes may be significantly higher than 3.1 (as shown in Table 5.2), which will slow the migration of these compounds, thereby increasing their susceptibility to biodegradation.
- 5. The availability of electron acceptors at this site does not appear to be limiting. In fact, the compost pile appears to have been a source of nitrate and sulfate to the groundwater (Figures 4.4 and 4.6). Although sulfate reduction and methanogenesis were not included as possible electron acceptors at this site, the compost pile made

- interpretation of the data difficult and inconclusive. Both sulfate reduction and methanogenesis may be occurring at the site.
- 6. Calibrated source concentrations in the models were somewhat higher than observed concentrations upgradient and crossgradient from the plume. This introduction of extra contaminant mass likely results in conservative predictions because additional BTEX mass must be biodegraded or discharged to the channel to produce the observed results.

The model simulation results demonstrate the potential for RNA to degrade the BTEX plume at the site without engineered source removal or treatment of the impacted medias. The patterns of degradation of the plume are feasible given the observed BTEX concentrations, the conservative assumptions made in constructing the simulation, and the strong evidence of biodegradation (Section 4).

SECTION 6

COMPARATIVE ANALYSIS OF REMEDIAL ALTERNATIVES

This section presents the development and comparative analysis of groundwater remedial alternatives for the UST Site at Beale AFB. The intent of this evaluation is to determine if RNA of groundwater is an appropriate and cost-effective remedial approach to consider when developing final remedial strategies for the site.

Section 6.1 presents the criteria used to evaluate groundwater remedial alternatives. Section 6.2 discusses the development and comparative analysis of remedial alternatives considered as part of this demonstration project. Section 6.3 provides a description of the selected remedial alternative(s). Section 6.4 provides a more detailed evaluation of the selected remedial alternative(s) using the defined remedial alternative evaluation criteria.

6.1 REMEDIAL ALTERNATIVE EVALUATION CRITERIA

The evaluation criteria used to identify appropriate remedial alternatives for shallow groundwater contamination at the site were adapted from those recommended by the USEPA (1988) for selecting remedies for Superfund sites [Office of Solid Waste and Emergency Response (OSWER) Directive 9355.3-01]. These criteria included (1) long-term effectiveness and permanence, (2) technical and administrative implementability, and (3) relative cost. The following sections briefly describe the scope and purpose of each evaluation criterion. This report focuses on the potential use of RNA to reduce BTEX concentrations in the shallow groundwater to concentrations that meet regulatory action levels.

6.1.1 Long-Term Effectiveness and Permanence

Each remedial technology or remedial alternative (which can be a combination of remedial approaches and technologies such as RNA and institutional controls) was analyzed to determine how effectively it will minimize groundwater plume expansion so that groundwater quality objectives can be achieved at a POC. The expected technical effectiveness based on case histories from other sites with similar conditions also is evaluated. The ability to minimize potential impacts to surrounding facilities and operations is considered. Also, the ability of each remedial alternative to protect both current and potential future receptors from potential risks associated with potential exposure pathways is qualitatively assessed. This evaluation criterion also included permanence and the ability to reduce contaminant mass, toxicity, and volume. Time to implementation and time until protection is achieved are described. Long-term reliability for providing continued

protection, including an assessment of potential for failure of the technology and the potential threats resulting from such a failure, is also evaluated.

6.1.2 Implementability

The technical implementation of each remedial technology or remedial alternative was evaluated in terms of technical feasibility and availability. Potential shortcomings and difficulties in construction, operations, and monitoring are presented and weighed against perceived benefits. Requirements for any post-implementation site controls such as LTM and land use restrictions are described. Details on administrative feasibility in terms of the likelihood of public acceptance and the ability to obtain necessary approvals are discussed.

6.1.3 Cost

The total cost (adjusted to present worth) of each remedial alternative was estimated for relative comparison. An estimate of capital costs, and operating and post-implementation costs for site monitoring and controls is included. An annual adjustment factor of 7 percent was assumed in present worth calculations. The annual adjustment factor is the difference between the rate of inflation and the cost of money (USEPA, 1993).

6.2 FACTORS INFLUENCING ALTERNATIVES DEVELOPMENT

Several factors were considered during the identification and screening of remedial technologies for addressing shallow groundwater contamination at the site. Factors considered included the objectives of the AFCEE natural attenuation demonstration program; contaminant, groundwater, and soil properties; current and future land uses; and potential receptors and exposure pathways. The following section briefly describes each of these factors and how they were used to narrow the list of potentially applicable remedial technologies to the final remedial alternatives considered for the UST Site.

6.2.1 Program Objectives

The intent of the RNA demonstration program sponsored by AFCEE is to develop a systematic process for scientifically investigating and documenting natural subsurface attenuation processes that can be factored into overall site remediation plans. The objective of this program and the specific demonstration at the UST Site at Beale AFB is to provide solid evidence of RNA of dissolved fuel hydrocarbons so that this information can be used to develop an effective groundwater remediation strategy. A secondary goal of this multi-site initiative is to provide a series of regional case studies that demonstrate that natural processes of contaminant degradation can often reduce contaminant concentrations in groundwater to below acceptable cleanup standards before completion of potential receptor exposure pathways.

Because the objective of this program is to study natural processes in the saturated zone rather than all contaminated media (e.g., soil, soil gas, etc.), technologies have been evaluated based primarily on their potential impact on shallow groundwater and phreatic soils. Technologies that can reduce vadose zone contamination and partitioning of contaminants into groundwater also have been evaluated. Many of the source removal

technologies evaluated in this section also will reduce soil and soil gas contamination, but it is important to emphasize that the remedial alternatives developed in this document are not intended to remediate all contaminated media. Additional program objectives set forth by AFCEE include cost effectiveness and minimization of waste. Applicable technologies that may meet these criteria include institutional controls and RNA. Soil excavation, slurry walls, sheet piling, groundwater pump and treat, air sparging, carbon adsorption, ex situ biological or chemical treatment, and onsite/offsite disposal are not considered attractive technologies for this site.

6.2.2 Contaminant Properties

The site-related contaminants considered as part of this demonstration at the site are the BTEX compounds. The primary source of this contamination is petroleum (gasoline and possibly other fuels) leaked or spilled during past aboveground and underground fuel storage at the site. Residual contamination at the site is present in the capillary fringe and possibly in saturated soils. The physiochemical characteristics of the individual BTEX compounds will greatly influence the effectiveness and selection of a remedial technology.

Gasoline is classified as an LNAPL with a liquid density of approximately 0.73 g/cm³ at 20°C (BEIA, 1989). Because gasoline is less dense than water, the LNAPL may become concentrated in the capillary fringe. Some of the individual gasoline constituents can either sorb to the soil matrix, dissolve into groundwater, or volatilize into soil vapor. Constituents in gasoline range from slightly to highly soluble in water. Overall solubility is approximately 300 mg/L. Gasoline is also a primary substrate for biological metabolism.

The BTEX compounds are generally volatile, highly soluble in water, and adsorb less strongly to soil than other hydrocarbons in a petroleum mixture. These characteristics allow the BTEX compounds to leach more rapidly from contaminated soil into groundwater, and to migrate as dissolved contamination (Lyman *et al.*, 1992). All of the BTEX compounds are highly amenable to *in situ* degradation by both biotic and abiotic mechanisms.

Benzene is very volatile, with a vapor pressure of 76 millimeters of mercury (mm Hg) at 20°C and a Henry's Law Constant of approximately 0.0054 atmosphere-cubic meters per mole (atm-m³/mol) at 25°C (Hine and Mookerjee, 1975; Jury *et al.*, 1984). The solubility of pure benzene in water at 20°C has been reported to be 1,780 mg/L (Verschueren, 1983). Benzene is normally biodegraded to carbon dioxide, with catechol as a short-lived intermediate (Hopper, 1978; Ribbons and Eaton, 1992).

Toluene is also volatile, with a vapor pressure of 22 mm Hg at 20°C and a Henry's Law Constant of about 0.0067 atm-m³/mol at 25°C (Pankow and Rosen, 1988; Hine and Mookerjee, 1975). Toluene sorbs more readily to soil media relative to benzene, but still is very mobile. The solubility of pure toluene in water at 20°C is approximately 515 mg/L at 20°C (Verschueren, 1983). Toluene has been shown to degrade to pyruvate, caetaldehyde, and completely to carbon dioxide via the intermediate catechol (Hopper, 1978; Wilson *et al.*, 1986; Ribbons and Eaton, 1992).

Ethylbenzene has a vapor pressure of 7 mm Hg at 20°C and a Henry's Law Constant of 0.0066 atm-m³/mol (Pankow and Rosen, 1988; Valsaraj, 1988). Ethylbenzene sorbs more

strongly to soils than benzene and toluene (Kenaga and Goring, 1980; Means *et al.*, 1980; Hassett *et al.*, 1983; Fetter, 1993). Pure ethylbenzene is also less soluble than benzene and toluene in water at 152 mg/L at 20°C (Verschueren, 1983; Miller *et al.*, 1985). Ethylbenzene ultimately degrades to carbon dioxide via its intermediate 3-ethylcatechol (Hopper, 1978; Ribbons and Eaton, 1992).

The three isomers of xylene have vapor pressures ranging from 7 to 9 mm Hg at 20°C and Henry's Law Constants of between 0.005 and 0.007 atm-m³/mol at 25°C (Mackay and Wolkoff, 1973; Hine and Mookerjee, 1975; Pankow and Rosen, 1988). A compilation of literature values for sorption coefficients suggests that xylenes sorb to soil with approximately the same strength as ethylbenzene (Wiedemeier *et al.*, 1995). Pure xylenes have water solubilities of 152 to 160 mg/L at 20°C (Bohon and Claussen, 1951; Mackay and Shiu, 1981; Isnard and Lambert, 1988). Xylenes can degrade to carbon dioxide via pyruvate carbonyl intermediates (Hopper, 1978; Ribbons and Eaton, 1992).

On the basis of these physiochemical characteristics, RNA, bioventing, soil vapor extraction, air sparging, biosparging, groundwater extraction, and air stripping technologies could all be effective options for collecting, destroying, and/or treating BTEX contaminants at the site. Some of these options are considered less desirable, however, after considering site-specific conditions.

6.2.3 Site-Specific Conditions

Three general categories of site-specific characteristics were considered when identifying remedial approaches for comparative evaluation as part of this demonstration. The first category was physical characteristics such as groundwater depth, hydraulic conductivity, gradient, flow direction, and soil type. The second category was the site geochemistry, or how the site contaminants are interacting with electron acceptors, microorganisms, and other site contaminants. Both of these categories influence the types of remedial technologies most appropriate for the site. The third category involved assumptions about future land use and potential receptor exposure pathways. Each of these site-specific characteristics have influenced the development of remedial alternatives included in the comparative evaluation.

6.2.3.1 Physical Characteristics

Site geology and hydrogeology have a profound effect on the transport of contaminants and the effectiveness and scope of required remedial technologies at a given site. Hydraulic conductivity is perhaps the most important aquifer parameter governing groundwater flow and contaminant transport in the subsurface. The velocity of the groundwater and dissolved contamination is directly related to the hydraulic conductivity of the saturated zone. Estimated hydraulic conductivity values from the slug tests performed at MW-1 ranged from 5 to 8 ft/day, characteristic of sand or dirty sand (Freeze and Cherry, 1979).

Zones of lower conductivity at the site were inferred from the slow recharge of water into some of the monitoring points (MP-12, MP-15, MP-17, and MP-19). These zones were modeled with a hydraulic conductivity approximately one order of magnitude less than the values obtained from aquifer tests at MW-1 (Section 5.4.1, Appendix D). The location of these lower conductivity zones and the hydraulic connection between the shallow

groundwater and the large drainage channel east-southeast of the site have created a flow pathway that runs from the source area toward MP-13 and MP-14 (Figures 3.10 and 4.1).

Site geology and hydrogeology also impact the types of practical engineered remedial technologies. For example, engineered solutions to plume containment are simplified because the preferential flow pathway already provides partial plume containment. Furthermore, less expense and time should be required to capture and treat the contaminant plume using extraction wells in areas of higher hydraulic conductivity which characterize typical preferential flow pathways. The drainage channel also acts as a natural interception trench for groundwater. In addition, the effectiveness of biosparging may increase as hydraulic conductivity increases because of reduced entry pressures and an increased radius of influence. Contaminant recovery may also be maximized when contaminants are not significantly sorbed to and retarded by phreatic soil.

6.2.3.2 Geochemical Characteristics

To satisfy the requirements of indigenous microbial activity and intrinsic bioremediation, the aquifer must also provide an adequate and available carbon or energy source (e.g., the contamination); electron acceptors; essential nutrients; and proper ranges of pH, temperature, and redox potential. Data collected as part of the fieldwork phase of this demonstration project and described in Sections 3 and 4 of this TS indicate that this site is characterized by adequate and available carbon/energy sources and electron acceptors to support measurable biodegradation of fuel hydrocarbon contamination by indigenous microorganisms. DO, ferric iron, and nitrate represent sources of electron acceptor capacity for the biodegradation of BTEX compounds at the site.

The average pH in shallow site groundwater is 6.8 and is within the optimal range for biodegradation of 6 to 8 standard pH units (Wiedemeier *et al.*, 1995). Fuel hydrocarbon-degrading microorganisms, however, have been known to thrive under a wide range of pH conditions (Chapelle, 1993). Groundwater data presented in Section 4 strongly support the conclusion that anaerobic biodegradation of BTEX is occurring at the site given the current geochemical conditions. Aerobic biodegradation should occur in the future with the removal of the compost pile.

Fuel-hydrocarbon-degrading microorganisms are ubiquitous, and as many as 28 hydrocarbon-degrading isolates (bacteria and fungi) have been discovered in different soil environments (Davies and Westlake, 1979; Jones and Eddington, 1968). Indigenous microorganisms have a distinct advantage over microorganisms injected into the subsurface to enhance biodegradation because indigenous microorganisms are well adapted to the physical and chemical conditions of the subsurface in which they reside (Goldstein *et al.*, 1985). Microbe addition was not considered a viable remedial technology for this site.

6.2.3.3 Potential Receptor Exposure Pathways

An exposure pathways analysis identifies the potential human and ecological receptors that could come into contact with site-related contamination and the pathways through which these receptors might be exposed. To have a complete exposure pathway, there must be a source of contamination, a potential mechanism(s) of release, a pathway of transport to an

exposure point, an exposure point, and a receptor. If any of these elements do not exist, the exposure pathway is considered incomplete and receptors will not come into contact with site-related contamination.

Evaluation of the potential long-term effectiveness of any remedial technology or remedial alternative as part of this demonstration project includes determining the potential for pathway completion. If a completed exposure pathway exists (e.g., surface water contact), potential long-term remedial options may still be sufficient to maintain exposure concentrations below regulatory action levels. Establishing site-specific, risk-based cleanup levels is beyond the scope of this TS.

Assumptions about current and future land uses at a site form the basis for identifying potential receptors, potential exposure pathways, reasonable exposure scenarios, and appropriate remediation goals. USEPA (1991) advises that the land use associated with the highest (most conservative) potential level of exposure and risk that can reasonably be expected to occur should be used to guide the identification of potential exposure pathways and to determine the level to which the site must be remediated.

The State of California can impose stricter guidelines where deemed appropriate. At Beale AFB, both groundwater and surface water themselves are considered potential receptors because of their potential use as drinking water sources and in accordance with state non-degradation policies. Therefore, any recommended remedial actions should incorporate a timeframe for reaching appropriate remedial goals for groundwater and surface water throughout the site.

The contaminant source areas at the site consist of soils containing residual LNAPL immediately south and west of the former excavation pit. Shallow groundwater is expected to serve as the predominant release and transport mechanism. The majority of shallow groundwater at the site discharges to the large drainage channel east-southeast of the site. No discharge of fuel-contaminated groundwater into the channel has been observed. Access to the site is not restricted for Base personnel, workers, or on-Base residents. Public access to the site as well as the Base is restricted.

Although it is not within the scope of this demonstration, a risk assessment may be required to evaluate potential risks. It is possible that both human and ecological receptors could come into contact with fuel hydrocarbon contamination that discharges from the shallow groundwater into the surface water. However, interpretation of surface water analytical data, contaminant distribution maps, groundwater gradient maps, and the modeling mass balance estimates suggests that currently only a very small amount of dissolved BTEX migrates into the drainage channel. Therefore, the probability for exposure to human and ecological receptors is very low.

Assumptions about hypothetical future land, groundwater, and surface water uses also must be made to ensure that the remedial alternative considered for shallow groundwater at the site is adequate and sufficient to provide long-term protection. No changes in land use are anticipated in the foreseeable future, so the current land use assumptions are appropriate for future scenarios. Current groundwater regulatory policy of the RWQCB requires that all

groundwater at the site be considered a potential drinking water source. Therefore, the groundwater itself must be considered a potential receptor.

In summary, the use of the RNA technology at this site will require that restrictions on shallow groundwater use be enforced until BTEX and TPH concentrations are reduced below California MCLs and taste and odor thresholds throughout the site. Modeling results indicate that RNA technology can achieve these remedial goals for all site groundwater within a reasonable time period. Contractors and government workers must be informed of the existence of the BTEX and TPH plume and a suitable Health and Safety Plan should be in place to ensure onsite worker safety. Site workers also must be made aware of the proper storage and disposal procedures for contaminated groundwater and restriction on its use.

These controls would be required until any remedial activities and natural attenuation reduce contaminants to levels that pose no risk. Should LTM or any future sampling efforts demonstrate that the contamination is impacting the surface water or has moved across the eastern drainage channel, it may be necessary to re-evaluate the remedial alternatives selected. If source reduction technologies such as bioventing or biosparging are implemented, they will have some impact on the short- and long-term land use options and some level of institutional control and worker protection during remediation will be required.

6.2.3.4 Remediation Goals for Shallow Groundwater

Model results suggest that BTEX and TPH compounds in shallow groundwater will naturally attenuate via biodegradation and that current discharges of contaminated groundwater to surface water are not significant. The selected remedial alternative(s) must be able to achieve groundwater concentrations protective of human health and the environment. The ultimate remedial objective for shallow groundwater is attainment of State of California surface water and groundwater quality objectives. California MCLs and taste and odor thresholds and federal MCLs are listed in Table 6.1 for dissolved fuel contaminants. Although it is unlikely that groundwater would be ingested by humans, California MCLs and taste and odor thresholds are currently the level of long-term protection required by The Water Quality Control Plan (Basin Plan) of the California Regional Water Quality Control Board (RWQCB), Central Valley Region (California RWQCB, 1994).

In summary, available data suggest that completed exposure pathways involving human and ecological receptors exposed to surface water may exist under current conditions. However, the probability for exposure of human and ecological receptors to significant contaminant concentrations is very low. A risk assessment for the surface water pathway may be required to evaluate risk from this pathway if future sampling indicates that BTEX contamination is reaching the surface water at detectable concentrations.

The data also suggest that there is no completed potential exposure pathway involving shallow groundwater, provided that use of shallow groundwater as a potable or industrial source of water is prohibited by institutional controls within approximately 500 feet of the UST Site. Thus, institutional controls are likely to be a necessary component of any groundwater remediation strategy for this site. The required duration of these institutional

controls may vary depending on the effectiveness of the selected remedial technology at reducing contaminant mass and concentrations in the groundwater.

TABLE 6.1

WATER QUALITY OBJECTIVES INTRINSIC REMEDIATION TS UST SITES 10-494, 10-495, AND 10-496 BEALE AFB, CALIFORNIA

Compound	Federal MCLs ¹ (µg/L)	California MCLs ² (μg/L)	Taste & Odor Threshold ³ (μg/L)
Benzene	5	1	170
Toluene	1,000	150	42
Ethylbenzene	700	700	29
Xylenes (Total)	10,000	1,750	17
Total Petroleum Hydrocarbons as Gasoline (TPH-g)	no standard	no standard	5

Drinking Water Regulations and Health Advisories, Office of Water, USEPA, May 1995.

6.2.4 Summary of Remedial Option Screening and Analysis

Several remedial options have been identified for use in treating the shallow groundwater at the site. Among the initial remedial technologies and approaches considered as part of this demonstration were institutional controls, LTM, groundwater extraction/containment via vertical wells and/or interceptor trenches; air sparging; physical containment using slurry walls, grout curtains, or sheet piles; funnel and reactive gate walls; and RNA. Screening was conducted by considering the program objectives of the AFCEE RNA demonstration, physiochemical properties of the BTEX compounds, and other site-specific characteristics such as hydrogeology, geochemistry, land use assumptions, potential receptor exposure pathways, and appropriate remediation goals. All of these factors will influence the technical effectiveness, implementation, and relative cost of technologies for remediating shallow groundwater underlying and migrating from the site.

Based on the screening process (Table 6.2), a single remedial approach — RNA with LTM and institutional controls — was retained for development of the remedial alternative. The

² California Code of Regulations (CCR), Title 22, Division 4, Chapter 15, Article 5.5, Section 64444, Maximum Contaminant Levels - Organic Chemicals.

³ Cal/EPA, 1998.

TABLE 6.2

TECHNOLOGIES AND PROCESS OPTIONS FOR GROUNDWATER REMEDIATION INITIAL TECHNICAL IMPLEMENTABILITY SCREENING OF

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation Treatability Study Beale AFB, California

General	Technology				Relative	
Response Action	Type	Process Option	Implementability	Effectiveness	Cost	Retain
Long-Term	Periodic	Confirmation	Sufficient space exists between the source and within	Necessary for all	Low	Yes
Monitoring	Groundwater/ Surface Water	Wells	the plume for an additional well to confirm the progress of remediation.	remediation strategies		
	Monitoring					
		Point-of-	A sufficient location exists downgradient of the plume	Necessary	Low	Yes
		Compliance	at the drainage channel to perform surface water sampling. This location will need to be monitored to			
			confirm that it is not impacted in the future.			
Institutional	Groundwater	Land Use	The plume lies within the base boundary, and land	Necessary	Low	Yes
Controls	Use Control	Control/Regulate	and groundwater use are under base jurisdiction.			
		well reimilis				
		Seal/Abandon	No production wells are known to exist in the current	Not required at this	Low	No
		Existing Wells	or predicted plume area.	site		
		Point-of-Use	No shallow groundwater is extracted from the plume	Poor	Moderate	No
		Treatment	area for any use.			
	Public	Meetings/	Base public relations and environmental management	Necessary	Low	Yes
	Education	Newsletters	offices have many information avenues to workers			
			and residents.			
Containment of	Hydraulic	Interceptor Trench	No likely receptors downgradient of site. Treatment	Low	Moderate	No
Plume	Controls	Collection	of extracted water would probably be necessary.		to High	
		Minimum	No likely receptors downgradient of site. Treatment	Low	High	No
		Pumping/ Gradient Control	of extracted water would probably be necessary.			
	Physical Controls	Slurry Walls/Grout	Limited effectiveness and costly.	Low	High	No
)	Sheet Piling	Limited effectiveness and costly	Low	High	SZ
					1	
	Keactive/Semi-	Biologically Active	Degration of BTEX can be stimulated by allowing	Moderate	High	°Z
	Permeable	Souo7	groundwater to flow through a nutrient-rich barrier.			
	Barriers		Although implementable, the technology is new and			
			unproveii.			

Table 6.2 (continued)

Technology Type	Process Option	Implementability	Effectiveness	Relative Cost	Retain
Biological	Oxygen and/or Nutrient Enhanced Biodegradation (Biosparging)	Nutrients/Oxygen are injected downgradient of plume to limit plume migration by enhancing biodegradation and reducing BTEX concentrations. Although implementable, the technology is not proven to be more effective than intrinsic remediation. In addition, DO is likely to be recharged naturally upgradient of plume at sufficent quantities in the future.	Moderate	Low	°Z
Chemical/ Physical	Intrinsic Remediation	A combination of natural biological, chemical, and physical removal mechanisms which occur to varying degrees on every site. Groundwater sampling at UST Site indicates that this is an ongoing remediation process.	High	Low	Yes
	Air Sparging (Volatilization)	Injection of air into contaminated aquifer creating a mass transfer of BTEX into air bubbles and vadose zone. May be difficult to implement due to near surface, low permeability layer.	Low	Moderate	N _o
Groundwater Extraction	Vertical Pumping Wells	Groundwater is pumped by installing submersible pumps in wells within or downgradient of plume. Could reduce the length of time required for natural attenuation processes to complete the groundwater restoration.	Moderate	Moderate	No
Biological	Bioreactors	High flow rates require excessive retention times and large reactors. BTEX is often volatilized in these systems.	Moderate	High	S _O
Chemical/ Physical	Air Stripping	Cost-effective technology for removing BTEX from groundwater at high flow rates. Difficult to permit.	High	Moderate	No
	Activated Carbon	Implementable at the low BTEX concentrations at the UST Site, but requires disposal of carbon.	Moderate	High (O&M)	No
	UV/Ozone Reactors	High flow rates require excessive retention times and large, expensive reactors.	Moderate	High	No
Direct Discharge	Direct Discharge to Industrial Waste Water Treatment Plant (IWWTP)	Implementable option only when an IWWTP is readily available and capable of handling BTEX, TPH, and hydraulic loading. IWWTP not available for this site	High	Low	No

Table 6.2 (continued)

General	Technology				Relative	
Response Action	Type	Process Option	Implementability	Effectiveness	Cost	Retain
Disposal	Sanitary Sewer	Sanitary Sewer	Implementable option when access to a sanitary sewer exists and loading is acceptable. Loading likely not acceptable to Base water treatment plant.	High	Low	No
	Treated Groundwater Reinjection	Vertical Injection Wells	Injection wells subject to clogging, high maintenance, and will be difficult to permit; aquifer is confined and/or semi-confined.	Moderate	Moderate	No
	•	Injection Trenches	Requires large trenches and will be difficult to permit.	Moderate	Moderate	No
	Discharge to Surface Waters		Generally requires NPDES or other discharge permit; not a viable option based on discussions with RWQCB.	High	Low	No
Source	Excavation/	Landfilling	Implementable at existing Base soils treatment facility	Moderate	High	No
Removal/Soil	Treatment		but excavation is expensive and gravelly soils may pose problem for treatment facility. Dewatering and treatment/disposal of groundwater might be required.			
Remediation		Biological Landfarming	Implementable at existing Base soils treatment facility but excavation is expensive and gravelly soils may pose problem for treatment facility.	Moderate	Moderate	No
		Thermal Desorption	More costly than existing Base soils treatment facility.	Moderate	High	No
	In Situ	Bioventing	Air injection/extraction to increase soil oxygen levels and stimulate biodegradation of fuel residuals. Limited effectiveness at UST Site due to near surface low permeability layer and shallow contamination and groundwater.	Low to High	Low	oN
		Soil Vapor Extraction	Limited effectiveness at UST Site due to near surface low permeability layer and shallow contamination and groundwater.	High	Low to Moderate	SZ _
		Soil Washing	Water and/or surfactant solution are forced through zones of residual contamination to enhance contaminant partitioning into the groundwater. Most effective in homogeneous sandy soils. May cause increased concentrations to discharge to surface water.	Low	High	NO

screening process provided sufficient support to select the optimal remedial alternative based on the available site characterization data; thus, a detailed comparative analysis was unnecessary. However, should additional site characterization activities or LTM indicate that an RNA strategy is not protective of surface water or that the contaminant concentrations in groundwater are not decreasing at the rates expected by the modeling results presented in Section 5, then a more detailed feasibility study should be performed.

6.3 DESCRIPTION OF THE SELECTED REMEDIAL ALTERNATIVE — REMEDIATION BY NATURAL ATTENUATION AND INSTITUTIONAL CONTROLS WITH LONG-TERM GROUNDWATER AND SURFACE WATER MONITORING

Remediation by natural attenuation (RNA) is achieved when natural attenuation mechanisms bring about a reduction in the total mass of a contaminant in the soil or dissolved in groundwater. RNA results from the integration of several subsurface attenuation mechanisms that are classified as either destructive or nondestructive. Destructive attenuation mechanisms include biodegradation, abiotic oxidation, and hydrolysis. Nondestructive attenuation mechanisms include sorption, dilution (caused by dispersion and infiltration), and volatilization. In some cases, RNA will reduce dissolved contaminant concentrations below numerical concentration goals intended to be protective of human health and the environment. As indicated by the evidence of RNA described in Section 4, these processes are occurring at the UST Site and will continue to reduce contaminant mass in the plume area.

Model BEALESW is intended to predict the fate and transport of dissolved BTEX compounds if engineered remedial action is not implemented at the site. To accomplish this, the model assumes a weathered source of BTEX in the shallow aquifer. Results suggest that the dissolved BTEX plume will decrease in concentration over time. The model conservatively predicts that the TPH and BTEX plume will be reduced to concentrations below groundwater quality objectives within approximately 15 years.

Implementation of RNA would require the use of institutional controls such as land use restrictions and well permit regulation until the groundwater quality objectives are attained. Land use restrictions may include placing long-term restrictions on soil excavation within the source area and recreational facilities (i.e., playgrounds). Restrictions on groundwater well installations within and downgradient from the plume area would also be required until groundwater quality objectives are achieved. The intent of these restrictions would be to reduce potential receptor exposure to contaminants by legally restricting activities within areas expected to be affected by site-related contamination. Since modeling results indicate that RNA could achieve groundwater quality objectives in a reasonable time period, institutional controls for groundwater would be temporary and would be expected to be removed within a reasonable timeframe. The combination of limited institutional controls, RNA, and a groundwater monitoring program is expected to be protective of future unrestricted groundwater uses.

LTM would be conducted annually as part of this remedial technology to evaluate the progress of natural attenuation processes. LTM includes two types of monitoring locations. Long-term monitoring wells are intended to determine if the behavior of the plume is changing and to monitor the progress of contaminant reductions at the site. POC monitoring locations are intended to detect contaminant movement outside the negotiated perimeter of containment and to trigger an action to manage the risk associated with such expansion if the contaminant concentrations at POC locations exceed the water quality objectives listed in Table 6.1.

As a part of the interim LTM plan for this site, groundwater and surface water monitoring would be performed at three groundwater monitoring wells and three surface water sampling locations (discussed in Section 7). Both groundwater monitoring wells and surface water sampling locations would be used as POC locations for this site. Groundwater monitoring wells would be used to evaluate changes in contaminant concentrations and ensure that the plume does not expand or migrate. Surface water sampling locations would be used to ensure that discharges into surface water do not exceed water quality objectives at any time. Both the groundwater and surface water POC locations will be used to determine if more active remediation methods are warranted.

As part of the interim LTM plan, monitoring well locations would be chosen to monitor upgradient groundwater geochemistry and the contaminant plume between the source area and the drainage channel at the location where the highest benzene and total BTEX concentrations were found during this study (MP-14). The surface water locations would be chosen in the large drainage channel. One location would be upgradient from the site, one location would be where the plume appears to discharge into the channel, and one location would be downgradient from the site. Additional details (including more detailed monitoring locations) for interim LTM of groundwater and surface water at the site are provided in Section 7.2. If implementation of the remedial alternative does not result in a decrease in dissolved contaminant concentrations within the predicted time frame or additional site characterization activities alter the site conceptual model presented in this report, additional corrective action may be necessary, and land use restrictions would require reevaluation.

Public education on the selected alternative should be developed to inform Base personnel and residents of the scientific principles underlying RNA. This education could be accomplished through debriefings, presentations, and posting of signs where appropriate. Periodic site reviews could also be conducted every year using data collected from the long-term groundwater monitoring program. The purpose of these periodic reviews would be to evaluate the extent of contamination, assess contaminant migration and attenuation through time, document the effectiveness of institutional controls at the site, and reevaluate the need for additional remedial actions at the site.

6.4 EVALUATION OF THE SELECTED REMEDIAL ALTERNATIVE — REMEDIATION BY NATURAL ATTENUATION AND INSTITUTIONAL CONTROLS WITH LONG-TERM GROUNDWATER AND SURFACE WATER MONITORING

6.4.1 Effectiveness

Section 5 of this document presents the results of the Bioplume II modeling of RNA at the UST Site. Model BEALESW assumes that site remediation relies entirely on natural attenuation mechanisms. Results of the model suggest that 33 percent of the dissolved BTEX can be removed from the groundwater system through natural attenuation, with the remaining 67 percent of the mass discharging to the drainage channel but not at measurable concentrations. Model results conservatively indicate that after 15 years, the dissolved BTEX concentrations in site groundwater would be less than or equal to $5 \,\mu\text{g/L}$. Based on the model results, the evidence of significant anaerobic biodegradation in the shallow groundwater (Section 4), and the potential for significant future aerobic biodegradation, RNA is an effective method to complete the restoration of site groundwater.

The effectiveness of this remedial alternative requires that excavations or drilling within the source area be conducted only by properly protected site workers. Long-term land use restrictions would be required to ensure that shallow groundwater is not pumped or removed for potable use within a radius of approximately 500 feet from the plume area. Existing health and safety plans should be enforced to reduce risks from additional excavation or from installing and monitoring additional wells.

Compliance with program goals is one component of the long-term effectiveness evaluation criterion. This alternative would satisfy program objectives designed to promote RNA as a component of site remediation and to scientifically document natural chemical attenuation processes. This alternative also satisfies program goals for cost effectiveness and waste minimization.

The RNA alternative is based on the effectiveness of natural processes that minimize contaminant migration and reduce contaminant mass over time and on the effectiveness of institutional controls. As described earlier, model results indicate that the BTEX plume will be contained and reduced within the source area via anaerobic biodegradation and future aerobic biodegradation. Therefore, the effectiveness of RNA with LTM and long-term institutional controls is favorable.

6.4.2 Implementability

RNA is not technically difficult to implement. Installation of monitoring wells and monitoring of groundwater and surface water are standard procedures. Long-term management efforts would be required to ensure proper sampling procedures are followed. Periodic site reviews should be conducted to confirm the adequacy and completeness of LTM data and verify the effectiveness of this remediation approach. There may also be administrative concerns associated with long-term enforcement of groundwater use restrictions. Future land use within the source area may be impacted by leaving contaminated soil and groundwater in place. Regulators would have to be informed of the

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benefits and limitations of the RNA option. Educational programs for Base personnel and residents are not difficult to implement. Where the effectiveness of this option has been supported, the initial regulatory reaction to this alternative has been positive.

6.4.3 Cost

The cost of RNA with the proposed interim LTM plan is summarized in Table 6.3. Capital costs are limited to the construction of two new monitoring wells. Included in the approximately \$104,000 total present worth cost estimate are the costs of maintaining institutional controls and long-term groundwater and surface water monitoring for a total of 15 years. Detailed cost calculations and assumptions are provided in Appendix F. Should additional site characterization activities or LTM indicate that an RNA strategy is not protective of surface water or that the contaminant concentrations in groundwater are not decreasing at the rates expected by the modeling results presented in Section 5, then a cost analysis comparison with more active remediation technologies should be performed.

TABLE 6.3

RNA WITH LTM ALTERNATIVE - COST ESTIMATE UST SITES 10-494, 10-495, AND 10-496 INTRINSIC REMEDIATION TS BEALE AFB, CALIFORNIA

<u>Capital Costs</u>	Cost
Design/Construct 2 LTM Wells	\$5,920
Monitoring Costs (per Sampling Event)	Cost per Event
Conduct Groundwater/Surface Water Sampling at 6 locations (per event) (quarterly for first year and annually thereafter for 15 years)	\$3,340
Maintain Institutional Controls/Public Education (15 years)	\$5,000
Project Management and Reporting (15 years)	\$1,100
Present Worth of Alternative 1 ^{a/}	\$103,880

a/ Based on an annual adjustment factor of 7 percent (USEPA, 1993).

Present worth costs and calculations are detailed in Appendix F.

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SECTION 7

LONG-TERM MONITORING PLAN

7.1 OVERVIEW

At the UST Site, LTM is a required component of the selected remedial alternative discussed in Section 6; therefore, a long-term groundwater monitoring plan must be developed. The purpose of the LTM component of the selected remedial alternative is to assess site conditions over time, confirm the effectiveness of the remedial alternative, assess compliance with regulatory cleanup goals, and evaluate the need for additional remediation. The LTM plan consists of identifying groundwater and surface water sampling locations and developing a sampling and analysis strategy to demonstrate attainment of site-specific RAOs. The strategy described in this section is designed to assess the effectiveness of the remedial alternative through measurement of the reduction of contaminant mass and the rate of groundwater remediation. In the event that data collected under this LTM program indicate that the implemented remedial alternative is insufficient to achieve regulatory levels considered to be protective of human health and the environment at the POC locations, additional engineered controls to augment the beneficial effects of RNA may be necessary.

7.2 GROUNDWATER MONITORING WELLS

An interim LTM plan is currently proposed for the UST Site based on the available site characterization data. This interim LTM plan would need to be updated if results from additional site characterization activities significantly change the current conceptual model for the site (e.g., if the groundwater plume is determined to have migrated east of the large drainage channel or if seasonal sampling of the surface water indicates that surface water contaminant concentrations are above water quality goals).

As part of the interim LTM plan, it is recommended that three groundwater monitoring wells be utilized at the site — MW-1 and two new wells (MW-3 and MW-4). The new wells should be fully screened across the shallow saturated zone between approximately 5 and 15 feet bgs to allow comparison with results from MW-1. The presence of free product/mobile LNAPL should be evaluated during installation of MW-3 and removed to the extent practicable in compliance with regulatory requirements. The proposed LTM well locations are shown on Figure 7.1. The location of MW-3 was chosen near MP-13 and MP-14, where the highest concentrations of BTEX and benzene were found during this investigation. The well is downgradient from the residual LNAPL source area along the preferential flowpath to the groundwater discharge point at the drainage channel. The location for MW-1 was chosen to continue to assess the upgradient electron acceptor conditions at the site. The location for

MW-4 was chosen to provide additional site characterization data on the east side of the large drainage channel, to establish hydraulic control for the site, and to verify plume stability.

7.3 SURFACE WATER SAMPLING LOCATIONS

Three surface water locations should be utilized at the site as part of the interim LTM. The proposed locations are shown on Figure 7.1 (SW-1 through SW-3). One location would be upgradient from the site, one location would be where the plume appears to discharge into the channel, and one location would be downgradient from the site.

The purpose of these locations is to verify that no contaminated surface water exceeding water quality objectives (Table 6.1) migrates from the site and beyond the area under institutional control.

7.4 GROUNDWATER AND SURFACE WATER SAMPLING

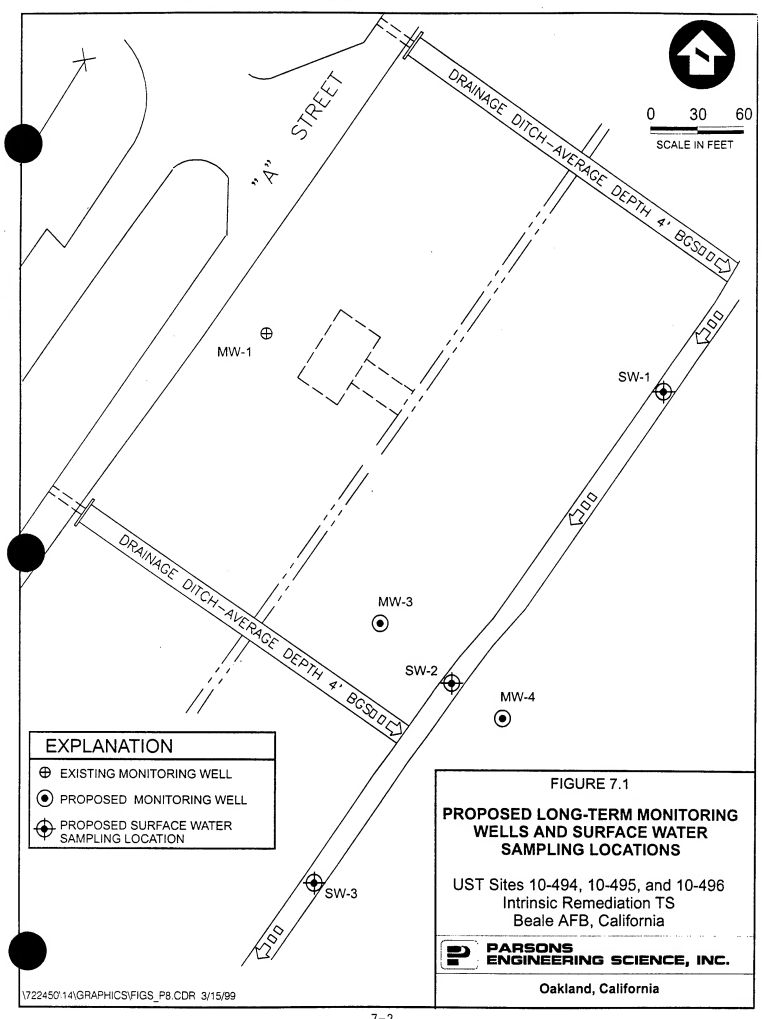
To ensure that sufficient contaminant removal is occurring to meet site-specific remediation goals, the long-term groundwater and surface water monitoring plan includes a comprehensive SAP. The groundwater and surface water locations should be monitored for BTEX and TPH-g to assure that the selected remedy is providing the anticipated level of risk reduction and remediation at the site. Results from the LTM locations will be used to verify that naturally occurring processes are effectively reducing contaminant mass and mobility. Reductions in toxicity will be implied by mass reduction and compliance with numerical RAOs.

7.4.1 Sampling Frequency

Because the results of this study are based on a single monitoring event, groundwater and surface water elevation measurements should be collected quarterly for at least one year to determine seasonal variations in groundwater flow direction and gradient at the site. Groundwater and surface water samples from the LTM wells and surface water locations also should be collected quarterly for the first year to determine if there are any significant seasonal variations in contaminant concentrations. Assuming there are no significant seasonal variations, sampling frequency should be reduced to annually after the first year. The LTM wells and surface water locations should be sampled annually until the plume disappears (estimated at no more than 15 years based on the modeling results discussed in Section 5). If the data collected during this time period support the effectiveness of the selected remedial alternative at this site (e.g., the plume disappears), it may be possible to reduce or eliminate sampling. If the data collected at any time during the monitoring period indicate the need for additional remedial activities at the site, sampling frequency should be adjusted accordingly.

7.4.2 Analytical Protocol

The proposed LTM wells should be sampled and analyzed to determine compliance with chemical-specific remediation goals and to verify the effectiveness of naturally-occurring remediation processes at the site. At the beginning of each sampling event, water levels and presence of free product/mobile LNAPL should be measured at all site monitoring wells and



in the surface water ditches. Groundwater samples collected from the LTM wells should be analyzed for the parameters listed in Table 7.1. Surface water samples should be analyzed for BTEX, TPH-g, DO, and temperature using the same analytical methods listed in Table 7.1. A site-specific groundwater SAP should be prepared prior to initiating the LTM program.

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TABLE 7.1

LONG-TERM MONITORING ANALYTICAL PROTOCOL FOR GROUNDWATER UST SITES 10-494, 10-495, AND 10-496 INTRINSIC REMEDIATION TS, BEALE AFB, CALIFORNIA

Field or Fixed-Base Laboratory	Field	Field	Field	Field	Field	Field
Sample Volume, Sample Container, Sample Preservation	Collect 100 mL of water in a glass container, acidify with hydrochloric acid per method	Collect 100 mL of water in a glass container	N/A	Collect 300 mL of water in biochemical oxygen demand bottles; analyze immediately; alternately, measure dissolved oxygen in situ	Collect 100–250 mL of water in a glass or plastic container; analyze immediately	Collect 100–250 mL of water in a glass container, filling container from bottom; analyze immediately
Recommended Frequency of Analysis	Quarterly for first year; annually thereafter for 15 years	Quarterly for first year, annually thereafter for 15 years	Quarterly for first year; annually thereafter for 15 years	Quarterly for first year; annually thereafter for 15 years	Quarterly for first year; annually thereafter for 15 years	Quarterly for first year; annually thereafter for 15 years
Data Use	May indicate an anacrobic degradation process due to depletion of oxygen, nitrate, and manganese	Same as above	Metabolism rates for microorganisms depend on temperature	The oxygen concentration is a data input to the Bioplume model; concentrations less than 1 mg/L generally indicate an anacrobic pathway	Aerobic and anacrobic processes are pH-sensitive	The redox potential of groundwater influences and is influenced by the nature of the biologically mediated degradation of contaminants; the redox potential of groundwater may range from more than 200 mV to less than -400 mV
Comments	Field only	Alternate method; field only	Field only	Refer to method A4500 for a comparable laboratory procedure	Protocols/ Handbook methods	Measurements are made with electrodes; results are displayed on a meter; samples should be protected from exposure to
Method/Reference	Colorimetrie A3500-Fc D	Colorimetric HACH 25140-25	E170.1	Dissolved oxygen meter	E150.1/SW9040, direct reading meter	A2580 B
Analysis	Ferrous (Fe ⁺²)	Ferrous (Fe ⁺²)	Temperature	Dissolved Oxygen	Hď	Redox potential

TABLE 7.1 (continued)

Analysis	Method/Reference	Comments	Data Use	Recommended Frequency of Analysis	Sample Volume, Sample Container, Sample Preservation	Field or Fixed-Base Laboratory
Conductivity	E120.1/SW9050, direct reading meter	Protocols/ Handbook methods	General water quality parameter used as a marker to verify that site samples are obtained from the same groundwater system	Quarterly for first year; annually thereafter for 15 years	Collect 100–250 mL of water in a glass or plastic container	Field
Nitrate (NO ₃ -1)	IC method E300 or method SW9056: colorimetric, method E353.2	Method E300 is a Handbook method; method SW9056 is equivalent procedure	Substrate for microbial respiration if oxygen is depleted	Quarterly for first year; annually thereafter for 15 years	Collect up to 40 mL of water in a glass or plastic container; cool to 4° C; analyze within 48 hours	Fixed-base
Aromatic hydrocarbons (BTEX)	SW8021	Handbook method; analysis may be extended to higher molecular weight alkyl benzenes	BTEX, are the primary target analyte for monitoring natural attenuation: BTEX concentrations must also be measured for regulatory compliance	Quarterly for first year; annually thereafter for 15 years	Collect water samples in a 40 mL VOA vial; cool to 4°C; add hydrochloric acid to pH 2	Fixed-base
Total Petroleum Hydrocarbons as gasoline (TPH-g)	SW8015B	Handbook method	Data used to monitor the reduction in concentrations of total fuel hydrocarbons (in addition to BTEX)	Quarterly for first year, annually thereafter for 15 years	Collect water samples in a 40 mL VOA vial; cool to 4°C; add hydrochloric acid to pH 2	Fixed-base

NOTES:

- "HACH" refers to the HACH Company catalog, 1990.
- "A" refers to Standard Methods for the Examination of Water and Wastewater, 18th edition, 1992.
- "E" refers to Methods for Chemical Analysis of Water and Wastes, U.S. Environmental Protection Agency, March 1979.
- "Protocols" refers to the AFCEE Environmental Chemistry Function Installation Restoration Program Analytical Protocols, 11 June 1992.
- "Handbook" refers to the AFCEE Handbook to Support the Installation Restoration Program (IRP) Remedial Investigations and Feasibility Studies (RI/FS), September 1993.
- "SW" refers to the Test Methods for Evaluating Solid Waste, Physical, and Chemical Methods, SW-846, U.S. Environmental Protection Agency, 3rd edition, Final Update III, December 1996.
- "ASTM" refers to the American Society for Testing and Materials, current edition.
- "RSKSOP" refers to Robert S. Kerr (Environmental Protection Agency Laboratory) Standard Operating Procedure. ∞ ¢.
 - "LUFT" refers to the state of California Leaking Underground Fuel Tank Field Manual, 1988 edition.

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SECTION 8

CONCLUSIONS AND RECOMMENDATIONS

This report presents the results of a TS conducted to evaluate the use of RNA with LTM for remediation of fuel-hydrocarbon-contaminated groundwater at the UST Site at Beale AFB. California. Specifically, the finite-difference groundwater model Bioplume II was used in conjunction with site-specific geologic, hydrologic, and laboratory analytical data to simulate the migration and biodegradation of fuel hydrocarbon compounds dissolved in groundwater. To collect the data necessary for the RNA demonstration, Parsons ES scientists collected soil, groundwater, and surface water samples from the site. Physical and chemical data collected under this program were supplemented (where necessary) with data collected during previous site characterization events.

Comparison of BTEX, TPH-g, electron acceptor, and biodegradation byproduct isopleth maps for the UST Site provides strong qualitative evidence of biodegradation of BTEX and TPH-g compounds. Geochemical data strongly suggest that biodegradation of fuel hydrocarbons is occurring at the site via the anaerobic processes of nitrate reduction and iron reduction. Aerobic biodegradation also is likely to occur in the near future due to recent changes in site conditions. Patterns observed in the distribution of hydrocarbons, electron acceptors, and biodegradation byproducts further indicate that biodegradation is reducing contaminant concentrations in site groundwater. Surface water data indicate that groundwater discharging to the drainage channel is not resulting in detectable surface water concentrations of these chemicals.

Site-specific geologic, hydrologic, and laboratory analytical data were used in the Bioplume II numerical groundwater model to simulate the effects of advection, dispersion, sorption, contaminant loading, and biodegradation on the fate and transport of the contaminant plume. Extensive site-specific data were used for model calibration and implementation. Model parameters that could not be obtained from existing site data were estimated using widely accepted literature values for soils similar to those found at the site. Conservative aquifer parameters were used to construct the Bioplume II model for this study, and, therefore, the model results presented herein may represent a worst-case scenario.

For the model simulation (BEALESW), it was assumed that weathering of the source areas would persist for the duration of the simulation. Model results conservatively indicate that after 15 years, the dissolved BTEX concentrations in groundwater would be less than or equal to $5\,\mu\text{g/L}$. Based on the model results, the evidence of significant anaerobic biodegradation in the shallow groundwater (Section 4), and the potential for significant future aerobic biodegradation, RNA with LTM is an effective method to complete the restoration of site groundwater.

The results of this study suggest that RNA of BTEX and TPH-g compounds is occurring at the site and intrinsic bioremediation should ensure protection of human health and the environment. Therefore, the Air Force recommends that RNA, LTM, and institutional controls be implemented at the site. Institutional controls such as restrictions on shallow groundwater use and protective equipment for onsite workers during future drilling or excavation activities would prevent completion of exposure pathways while site remediation is in progress.

To verify the results of the Bioplume II modeling effort and to ensure that the selected remediation is progressing at rates sufficient to meet objectives, groundwater from three LTM wells should be sampled and analyzed for the parameters listed in Table 7.1. The surface water locations should be analyzed for BTEX, TPH-g, DO, and temperature by the methods listed in Table 7.1.

Figure 7.1 shows suggested locations for the interim LTM groundwater monitoring wells for surface water sampling. The presence of free product/mobile LNAPL should be evaluated during installation of any new LTM well(s) and removed to the extent practicable in compliance with regulatory requirements. Because of the removal of the compost pile near the source area at this site, the groundwater geochemistry can be expected to change within the next few years. Therefore, it is recommended that the assimilative capacity of site groundwater be reevaluated during LTM.

The LTM plan presented in Section 7 of this report will serve as an interim LTM until installation of the LTM wells and completion of the first year of sampling. The interim LTM wells and surface water locations should be sampled quarterly for the first year. Thereafter, the LTM should be updated as necessary and sampling should be reduced to annually if no significant seasonal changes in groundwater flow direction and gradient and contaminant concentrations are measured. Sampling should continue until the BTEX and TPH-g concentrations fall below regulatory levels of concern (estimated at 10 to 15 years). Sampling could decrease in frequency if the plume disappears, as dictated by the analytical results. If any BTEX or TPH-g concentrations detected in surface water exceed the water quality objectives or concentrations in groundwater are increasing significantly greater than that predicted by modeling results, additional corrective actions and comparative cost analyses may be required to remediate groundwater at the site.

SECTION 9

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APPENDIX A

BOREHOLE LOGS, MONITORING POINT AND WELL CONSTRUCTION SUMMARY

MONITORING POINT AND WELL CONSTRUCTION SUMMARY

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

				Total	Screened	Sand Pack	Bentonite	Grout
	Date		Diameter	Depth	Interval	Interval ⁽²⁾	Interval	Interval
Location	Completed	Material	(inches)	(feet bgs)	(feet bgs)	(feet bgs)	(feet bgs)	(feet bgs)
MP-01	3/1/96	PVC	0.50	9.75	4.75 - 9.75	3.5 - 9.75	0.0 - 3.5	
MP-03	2/29/96	PVC	0.50	8.8	3.8 - 8.8	3.0 - 8.8	0.0 - 3.0	
MP-08	2/27/96	SS	0.50	17.0	16.5 - 17.0		0.0 - 16.0	
MP-09	2/27/96	PVC	0.50	10.5	5.0 - 10.0	3.0 - 10.0	0.0 - 3.0	
MP-10	2/28/96	PVC	0.50	10.5	4.0 - 9.0	3.0 - 10.5	0.0 - 3.0	
MP-11	2/28/96	PVC	0.50	10.5	4.0 - 9.0	3.0 - 9.0	0.0 - 3.0	
MP-12	2/27/96	PVC	0.50	11.5	4.8 - 9.8	3.0 - 11.5	0.0 - 3.0	
MP-13	2/27/96	PVC	0.50	10.5	5.0 - 10.0	3.0 - 10.5	0.0 - 3.0	
MP-14	2/27/96	PVC	0.50	10.0	4.7 - 9.7	3.0 - 9.7	0.0 - 3.0	
MP-15	2/27/96	PVC	0.50	13.0	8.0 - 13.0	5.0 - 13.0	0.0 - 5.0	
MP-16	2/29/96	SS	0.50	19.5	19.0 - 19.5		0.0 - 16.5	
MP-17	2/28/96	PVC	0.50	11.0	4.5 - 9.5	3.0 - 11.0	0.0 - 3.0	
MP-18	2/26/96	PVC	0.50	10.5	4.9 - 9.9	4.0 - 9.9	0.0 - 4.0	
MP-19	2/28/96	PVC	0.50	-13.0	7.8 - 12.8	4.0 - 12.8	0.0 - 4.0	
MP-20	2/28/96	PVC	0.50	10.5	4.8 - 9.6	3.0 - 9.6	0.0 - 3.0	
MP-21	2/27/96	PVC	0.50	10.5	4.5 - 9.5	3.0 - 10.5	0.0 - 3.0	
MP-22	2/27/96	PVC	0.50	10.5	5.0 - 10.0	4.0 - 10.0	0.0 - 4.0	~-
MP-23	3/1/96	SS	0.50	17.5	17.0 - 17.5		0.0 - 16.0	
MP-24	2/26/96	PVC	0.50	10.5	4.6 - 9.8	4.0 - 9.8	0.0 - 4.0	
MP-25	2/28/96	PVC	0.50	9.5	4.0 - 9.0	3.0 - 9.0	0.0 - 3.0	
MP-26	2/28/96	PVC	0.50	10.5	4.0 - 9.0	3.0 - 9.0	0.0 - 3.0	••
MP-27	2/28/96	PVC	0.50	10.5	5.0 - 10.0	4.0 - 10.0	0.0 - 4.0	
MW-01 (b)	2/21/95	PVC	2.0	16.5	4.5 - 14.5	3.5 - 16.5	1.5 - 3.5	0.5 - 1.5
MW-02 (b)	2/21/95	PVC	2.0	18.0	13.0 - 18.0	11.0 - 18.0	8.0 - 11.0	0.5 - 8.0

⁽a) For stainless steel sacrifical points, point is driven into native material (no sand pack).

PVC: Polyvinyl chloride

SS: Stainless steel

⁽b) Monitoring wells were installed by Metcalf & Eddy (1995).

PARSONS ENGINEERING SCIENCE, INC.-

BORING NUMBER: MP-01

PROJECT NUI	MBER: 722450.14020	PROJECT NAME:		
CLIENT: AF	CEE	Beale Intrinsic Remediation Treatability Study		
LOCATION:	Beale Air Force Base, California			
	UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe		
GEOLOGIST:	Mark Vessely	HOLE DIAMETER: 2 inches		
COMPLETION	DATE: 3/1/96	TOTAL DEPTH: 9.75 feet below ground surface		

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -	ND CM		Ξ.	JE ///	SC Fill?	Dark Brown Clayey SAND with fine to medium gravel [Fill?]
5 —	¥	BE-MP01-5 [poor recovery]	NR/80		ML	Red Brown Clayey SILT with some fine sand and gravel, moist, no hydrocarbon odors
_			NR/75		SM	Dark Brown Silty Medium SAND with gravel, moist, no hydrocarbon odors
10 -						Total Depth 9.75' bgs [no water in borehole at time of drilling]
15 —						
-			-			
20						
25 —						
-						

— - Contact.

- Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:		
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study		
LOCATION: Beale Air Force Base, California			
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe		
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches		
COMPLETION DATE: 2/29/96	TOTAL DEPTH: 10.5 feet below ground surface		

					,	
DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0					fill?	Dark Brown Clayey Sandy Fine GRAVEL, moist to wet, angular to rounded gravel, no hydrocarbon odors
- 1	T		0.8/0		ML/ CL	Red-Brown Gravelly Clayey SILT/Silty CLAY with fine sand, moist, to wet, sub-rounded gravel, abundant organics, no hydrocarbon odor, soft
5 -	室	BE-MP03-4.5	1.0/0		GC	Red-Brown Sandy Clayey Fine GRAVEL, moist to saturated, no hydrocarbon odors, angular gravel, fine to medium sand
-					GC	Mottled Green and Brown Silty Clayey GRAVEL, moist, no hydrocarbon odors
10 -			1.1/0		CL/ ML	Mottled Green and Brown Gravelly Silty CLAY or Clayey SILT with Sand, moist, no hydrocarbon odors
15 —						Total Depth 10.5' bgs

- Contact

- Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME: Beale Intrinsic Remediation Treatability Study		
CLIENT: AFCEE			
LOCATION: Beale Air Force Base, California			
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe		
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 1 inch		
COMPLETION DATE: 2/27/96	TOTAL DEPTH: 18 feet below ground surface		

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
10 —			0.0/0		ML/CL SM ML/CL SM	O - 10.5' bgs same as MP-09 Same as above (see MP-09) Brown Gravelly Silty Fine to Coarse SAND, moist to wet, no hydrocarbon odors Mortled Olive-Green and Brown Clayey SILT/Silty CLAY with sand, moist, stiff Brown Gravelly Silty Fine to Coarse SAND, moist to wet, no hydrocarbon odors Total Depth 18' bgs

Contact.

- Contact approximately located.

- First encountered groundwater.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/27/96	TOTAL DEPTH: 10.5 feet below ground surface

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
5	▼		5.9/40		fill	FILL: Sand, Silt, Gravel and Clay, moist Compost: Primarily wood chips, moist to wet, no hydrocarbon odors
_		BE-MP09-6	0.0/15	1111	ML	Brown Clayey SILT with sand and gravel, moist to wet, no hydrocarbon odors Brown Gravelly Silty Fine to Coarse SAND with clay, saturated, no
10		BE-MP09-10	0.0/0		SM ML/ CL	Brown Gravelly Silty Fine to Coarse SAND with clay, saturated, no hydrocarbon odors Mottled Olive-Green and Brown Clayey SILT/Silty CLAY, moist, stiff, black organics Total Depth 10.5' bgs
25 —						

Contact.

- Contact approximately located.

PID - Photoionization detector reading.

- First encountered groundwater level.

TVH - Total Volatile Hydrocarbon meter reading.

- Equilibrated groundwater level.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/28/96	TOTAL DEPTH: 10.5 feet below ground surface

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -					ML	Brown Clayey SILT with fine gravel and trace sand, moist to wet, soft, Color change to Black or Dark Grey at 1.5' bgs, hydrocarbon odor [heavy HC?]
5 —	¥	:	0.8/40		ML	Brown Gravelly Sandy SILT, moist to wet, stiff, some lenses of gravelly silty fine to medium sand, moderate gasoline odor at 5' bgs with color change to green.
-	*		20/60		SM/ SW	Brown Silty Gravelly Fine to Coarse SAND, saturated, slight hydrocarbon odor, rounded fine gravel
10 -			1.4/3		ML CL	Mottled Olive-Green and Brown Clayey SILT with sand and gravel, moist. no hydrocarbons odor, stiff Olive-Green Silty CLAY, very stiff, no hydrocarbon odor Total Depth 10.5' bgs
_		*				
15 —						
20			. *			
25 —						
				:		

- Contact.

- Contact approximately located.

PID - Photoionization detector reading.

- First encountered groundwater level.

TVH - Total Volatile Hydrocarbon meter reading.

- Equilibrated groundwater level.

PROJECT NUMBER: 722450.14020	PROJECT NAME:		
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study		
LOCATION: Beale Air Force Base, California			
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe		
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches		
COMPLETION DATE: 2/28/96	TOTAL DEPTH: 10.5 feet below ground surface		

					,	
DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -					ML	Red-Brown Clayey SILT, moist to wet, soft, no hydrocarbon odors
_	_		0.0/0		CL	Red-Brown Silty CLAY, moist, stiff, no hydrocarbon odors Color change at ~4' bgs to mottled olive-green and brown with abundant organics
5 -	¥		90/60		SM	Brown Silty Gravelly Fine to Coarse SAND, moist to wet, minor clay Slight gasoline odor at 7 to 7.5' bgs
-					sw	Brown Gravelly Medium to Coarse SAND with silt and clay, saturated, slight gasoline odor [?], sub-rounded gravel
10 -			0.0/0		CL	Mottled Grey and Brown Silty CLAY, moist, hard, no hydrocarbon odors
						Total Depth 10.5' bgs
15 —						
			•			÷
-						
20 —						
-			!			
25 —						
-						

- Contact.

- Contact approximately located.

PID - Photoionization detector reading.

- First encountered groundwater level.

TVH - Total Volatile Hydrocarbon meter reading.

- Equilibrated groundwater level.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/27/96	TOTAL DEPTH: 11.5 feet below ground surface

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -		-			ML	Brown Clayey SILT with sand and gravel, wet near surface, soft, below 1' bgs becomes dry to damp, stiff and friable, no hydrocarbon odors, abundant organics
-		BE-MP12-4	0.0/0		ML	Mottled Olive-Brown and Brown Clayey SILT, moist to damp, stiff, abundant organics, no hydrocarbon odor
5 -	¥¥				SM	Mottled Olive-Green and Brown Silty Very Fine to Fine SAND, wet to saturated, minor gravel, no hydrocarbon odor Becomes Brown at ~5.5' bgs
-			0.0/0	П	ML	Mottled Olive-Green and Brown Clayey SILT, moist, very stiff
-					ML	Mottled Olive-Green and Brown, Clayey Gravelly Sandy SILT, moist to wet, no hydrocarbon odors
10 -			0.0/0		CL	Mottled Olive-Green Silty CLAY with fine sand and fine gravel, moist, stiff, no hydrocarbon odor
20						Total Depth 11.5' bgs

Contact.

- Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:		
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study		
LOCATION: Beale Air Force Base, California			
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe		
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches		
COMPLETION DATE: 2/27/96	TOTAL DEPTH: 10.5 feet below ground surface		

БЕРТН (fect)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
5 —	▼		0.0/0		ML	Red-Brown Clayey SILT, moist, occasional fine gravel, no hydrocarbon odors, some rootlets, soft Becomes friable and dry at ~2' bgs At 3' bgs soils are gravelly and sandy
		BE-MP13-6.5	230/50		SM	Mottled Grey-Brown and Brown Silty Gravelly Fine to Coarse SAND, moist to wet, angular gravel, slight to moderate gasoline odor
10 -			210/65		ML/ CL	Mottled Olive-Green and Brown Clayey SILT/Silty CLAY, moist, slight to no hydrocarbon odors, very stiff to hard Total Depth 10.5' bgs
15 —			·			
20 —						
25 —						

Contact.

- Contact approximately located.

PID - Photoionization detector reading.

- First encountered groundwater level.

- Equilibrated groundwater level.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/27/96	TOTAL DEPTH: 10 feet below ground surface

DEPTH (feet)	GW ELEVATION	SAMPLE	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0					ML	Black Gravelly SILT with sand, wet, fine angular gravel
-	_		0.0/0		ML	Red-Brown Clayey Gravelly SILT with sand, moist, soft, no hydrocarbon odors
5 -		BE-MP14-7	475/390		SM	Red-Brown Gravelly Silty Fine to Medium SAND, moist, trace clay, no hydrocarbon odor Moderate gasoline odor at 6.5'
10 -			118/26		ML/ CL	Mottled Olive-Green and Brown Clayey SILT/Silty CLAY with fine sand, moist to wet, slight gasoline odor, stiff Total Depth 10' bgs
15 —			·	•		
25 —						·

- Contact.

- Contact approximately located.

PID - Photoionization detector reading.

- First encountered groundwater level.

TVH - Total Volatile Hydrocarbon meter reading.

- Equilibrated groundwater level.

PROJECT NUMBER: 722450.14020	PROJECT NAME:		
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study		
LOCATION: Beale Air Force Base, California	·		
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe		
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches		
COMPLETION DATE: 2/27/96	TOTAL DEPTH: 13 feet below ground surface		

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -			0.0/0		ML	Red-Brown Clayey SILT, wet, soft, rootlets, trace sand and fine gravel
5 -	T		0.0/0		ML	Red-Brown Gravelly SILT with Clay and Sand, moist, no hydrocarbon odor, fine angular gravel, fine sand Color change to Olive-Green at ~ 5.5' bgs
10 -		BE-MP15-9.5	0.0/0		SM	Olive-Green Gravelly Silty Fine to Medium SAND, wet, minor clay [?], no hydrocarbon odors, fine sub-angular to angular gravel, very silty Moderate gasoline odor at ~ 9' bgs
_			0.0/0		ML	Mottled Olive-Green and Brown Clayey SILT, moist to wet, hard, slight hydrocarbon odor [?] Color change to Olive Green at ~12' bgs
15 — - - - - 20 —			-			Total Depth 13' bgs [No water in borehole at time of drilling]
25 —						

Contact.

-- - Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PARSONS ENGINEERING SCIENCE, INC.—

BORING NUMBER: MP-16

PROJECT NUMBER: 722450.14020	PROJECT NAME:		
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study		
LOCATION: Beale Air Force Base, California			
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe		
GEOLOGIST: Mark Vessely	HOLE DIAMETER: 2 inches		
COMPLETION DATE: 2/29/96	TOTAL DEPTH: 19.5 feet below ground surface		

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
5 -						0 - 13' bgs same as MP-15 Driller reports softness at 11' bgs
15 —			0.0/0		ML	Olive-Green Clayey SILT, moist, no hydrocarbon odors
	革		0.2/10 1.0/10		SM	Red-Brown silty SAND with clay [borehole filling with water after sampling past 17.5]
-			1.0/10	Ш	ML	Red-Brown Gravelly SILT with coarse sand, wet, no hydrocarbon odor
25 -						Total Depth 19.5' bgs

Contact.

- Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/28/96	TOTAL DEPTH: 11 feet below ground surface

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -				\coprod	ML	Red-Brown Clayey SILT, moist to wet, rootlets, occasional fine Gravel and Sand, no hydrocarbon odors, soft
					CL	As above except Silty CLAY
5 -			0.0/0		GM/ ML	Red Brown Sandy Silty GRAVEL/Gravelly SILT, moist to wet, no hydrocarbon odors
_	¥ ¥		17/40		ML	Red Brown Clayey SILT with Sand and Gravel, moist to wet, no hydrocarbon odors
-					GM	Brown Silty Sandy GRAVEL, saturated, fine angular gravel, fine to coarse sand, no hydrocarbon odors
10 -			0.8/0		ML	Tan Clayey SILT, moist, no hydrocarbon odors
_				-1-1-		Total Depth 11' bgs
-						
15 —						
-					:	
_						
20 —						
_						
25 —						
-						
_						

- Contact.

- Contact approximately located.

PID - Photoionization detector reading.

First encountered groundwater level.

TVH - Total Volatile Hydrocarbon meter reading.

- Equilibrated groundwater level.

ocarbon meter

ng. NA - Not Applicable.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/26/96	TOTAL DEPTH: 10.5 feet below ground surface

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -	¥	BE-MP18-3.0	0.0/0		ML	Brown Clayey Gravelly sandy SILT, wet, soft, very fine to fine sand
5 -	¥				ML	Light Brown Sandy SILT, moist, very fine sand
_	_		0.0/0		SM	Mottled Light Brown Silty Fine SAND, moist to wet
10 -			0.0/0		GP	Mottled Brown and Grey-Green Sandy Fine GRAVEL with silt and clay, saturated, angular gravel, fine to coarse sand
-						Total Depth 10.5' bgs
				!		
-						
15 —						
			-			
-				·		
20 —						
-						
25 —						
-						

- · Contact.

Contact approximately located.

PID - Photoionization detector reading.

- First encountered groundwater level.

- Equilibrated groundwater level.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/28/96	TOTAL DEPTH: 13 feet below ground surface

TH et)	ELEVATION	SAMPLE	SOIL HEADSPACE PID/TVH (ppmv)	IC LOG	CLASS	
DEPTH (feet)	GW ELE	SAM	SC HEAD PID/ (PPI	GEOLOGIC LOG	SOIL C	GEOLOGIC DESCRIPTION
0 -		•	0.0/0		ML/ CL	Red-Brown Clayey SILT/Silty CLAY with fine Gravel, moist to saturated (saturated near surface), no hydrocarbon odor, soft
5 -	_	BE-MP19-7	0.0/0		GC	Red Brown Clayey Sandy Fine GRAVEL with Silt, moist to wet, no hydrocarbon odors, sand is fine to coarse, angular gravel
10 -			0.0/0		sc	Red Brown Clayey Gravelly Fine to Coarse SAND, moist to wet, no hydrocarbon odors
					CL GC	Mottled Olive-Green and Brown Sandy Gravelly CLAY with SILT, moist to wet, no hydrocarbon odors, hard Red-Brown and Grey Clayey Sandy Fine GRAVEL, damp to moist, no hydrocarbon odors
15			·	2.252		Total Depth 13' bgs [No water in hole at 13' bgs]
20 —						
25 —						
_						

Contact.

- Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/28/96	TOTAL DEPTH: 10.5 feet below ground surface

ДЕРТН (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -	¥¥		0.0/0		ML/ CL	Red-Brown Clayey SILT/Silty CLAY with fine Gravel, wet to saturated near surface, organics, soft Change at 1.5' bgs to damp and friable - very stiff, no hydrocarbon odor, more silt than clay
5 -			0.6/0		GC/ SC	Red Brown Clayey Sandy Fine GRAVEL/Clayey Gravelly Fine to Coarse SAND, saturated, no hydrocarbon odor Mottled Brown and Olive-Green, as above at 8' bgs
10 -			1.2/10		CL	Mottled Brown and Olive Green Silty CLAY, moist to wet, very stiff, no hydrocarbon odors Total Depth 10.5' bgs
15 —						
20 —			. •			·
25 —		·				·

- Contact.

-- - Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/27/96	TOTAL DEPTH: 10.5 feet below ground surface

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -	¥¥		0.0/0		ML	Red-Brown Clayey SILT, moist to wet, abundant organics, minor gravel, soft
_				П	ML	Red-Brown Gravelly Sandy SILT, wet to saturated
5 -			0.0/0		sw	Brown Gravelly Fine to Coarse SAND with silt, saturated, no hydrocarbon odors
-			0.00		GP	Brown Sandy Fine GRAVEL with silt, saturated, no hydrocarbon odor, fine to coarse sand
10 -			0.0/0		ML	Olive-Green Clayey Gravelly SILT with sand, moist, no hydrocarbon odors, stiff
20 —						Total Depth 10.5' bgs

- - Contact.

--- - Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/27/96	TOTAL DEPTH: 10.5 feet below ground surface

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -	_		0.0/0		ML	Red-Brown Clayey SILT with sand, moist to wet, soft, no hydrocarbon odors, rootlets
5 —	¥	BE-MP22-7	20/10		SM/ ML	Red-Brown Gravelly Silty Fine to Medium SAND/Sandy SILT with clay, moist to wet, fine gravel, some organics At 6.5' bgs, gasoline odors and greenish color, saturated at 7.5 - 8' bgs 3" layer of Silty CLAY at ~ 9.5-9.8' bgs No apparent hydrocarbon odors below 9' bgs
10			0.0/0	<u> </u>		Total Depth 10.5' bgs
15 —			o ⁻			
20						·
25 —						
-						

Contact.

- Contact approximately located.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

- Equilibrated groundwater level.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Michael Phelps	HOLE DIAMETER: 2 inches
COMPLETION DATE: 3/1/96	TOTAL DEPTH: 17.5 feet below ground surface

					,	
DЕРТН (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
5		BE-MP23-6.5 BE-MP43-7.0	0.7/10 1.1/70 1.7/70			0-16.5' same as MP-17
			1.00		ML	Olive-Green Clayey Gravelly SILT, moist, no hydrocarbon odors
			1.8/0		ML	Red-Brown Sandy SILT with gravel, moist, no hydrocarbon odors
20 —					ML	Total Depth 17.5' bgs

- Contact.

--- - Contact approximately located.

- First encountered groundwater level

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:				
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study				
LOCATION: Beale Air Force Base, California					
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe				
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches				
COMPLETION DATE: 2/26/96	TOTAL DEPTH: 10.5 feet below ground surface				

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0					ML	Brown SILT [TOPSOIL], wet, soft, abundant grasses
					GM	Brown Silty Fine to Medium GRAVEL with fine sand, angular, gravel, moist
-	BE-MP24-3 0.4/0		0.4/0		ML	Brown Clayey Sandy SILT, wet to saturated, very soft
5 -					SM	Brown Silty Gravelly Fine to Medium SAND, wet
-			0.1/0		GM	Brown Silty Fine GRAVEL with sand, moist to wet, angular gravel
-					SP	Mottled Brown Gravelly Fine SAND with silt, wet, fine angular gravel
10 -			0.0/0		ML/ CL	Mottled Grey and Brown Clayey SILT/ Silty CLAY, moist, stiff
-						Total Depth 10.5' bgs
15						

- Contact.

--- - Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:					
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study					
LOCATION: Beale Air Force Base, California						
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe					
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches					
COMPLETION DATE: 2/28/96	TOTAL DEPTH: 9.5 feet below ground surface					

ДЕРТН (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
0 -	T		0,2/0		ML	Brown to Red-Brown Clayey SILT, moist to wet, no hydrocarbon odors, soft, rootlets, trace of fine sub-rounded gravel
5 —	¥	BE-MP25-7	0.1/0		SM	Brown Gravelly Silty Fine to Coarse SAND, moist to saturated, angular gravel, no hydrocarbon odors, some lenses of Gravelly SAND (SW)
10 -			0.8/0		ML	Mottled Olive-Green and Brown Silty CLAY/Clayey SILT, moist to wet, stiff, no hydrocarbon odors Total Depth 9.5' bgs
15 —						
20 —						
25 —						
-						

- Contact.

- Contact approximately located.

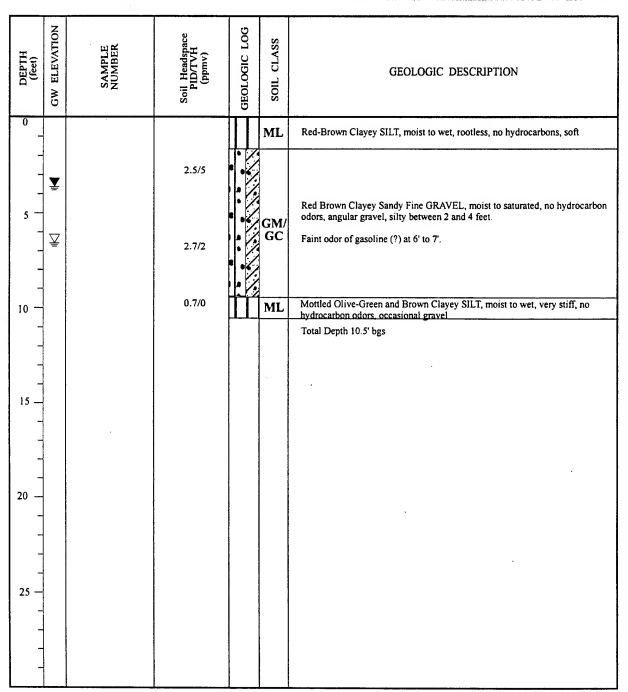
- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/28/96	TOTAL DEPTH: 10.5 feet below ground surface



- - Contact.

--- - Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

PROJECT NUMBER: 722450.14020	PROJECT NAME:
CLIENT: AFCEE	Beale Intrinsic Remediation Treatability Study
LOCATION: Beale Air Force Base, California	
UST Sites 10-494, 10-495 and 10-496	DRILLING METHOD: Geoprobe
GEOLOGIST: Marcus Pierce	HOLE DIAMETER: 2 inches
COMPLETION DATE: 2/28/96	TOTAL DEPTH: 10.5 feet below ground surface

DEPTH (feet)	GW ELEVATION	SAMPLE NUMBER	SOIL HEADSPACE PID/TVH (ppmv)	GEOLOGIC LOG	SOIL CLASS	GEOLOGIC DESCRIPTION
5 —	▼		0.3/0		ML	Red-Brown Clayey SILT with fine gravel and sand, angular to sub-angular gravel; moist to wet, rootlets, no hydrocarbon odors, stiff below 1' bgs Gravelly between 2.6 and 3.6' bgs Some thin lenses of Silty Sandy Fine Gravel Saturated at ~5 - 6' bgs, no hydrocarbon odors
10 -			0.3/5		CL/ ML	Mottled Olive-Green, Brown and Black Silty CLAY/Clayey SILT, moist to wet, no hydrocarbon odors, trace fine gravel, stiff Total Depth 10.5' bgs
15 —						
20 -						
25 —						

- - Contact.

- Contact approximately located.

- First encountered groundwater level.

- Equilibrated groundwater level.

PID - Photoionization detector reading.

TVH - Total Volatile Hydrocarbon meter reading.

APPENDIX B

SOIL AND GROUNDWATER ANALYTICAL RESULTS FROM PREVIOUS INVESTIGATIONS

UST SITE SUMMARY TANK NOS. 10-494, 10-495 AND 10-496

Tanks 10-494, 10-495, and 10-496 were historically located near the intersection of 30th and A Streets at Beale AFB. In accordance with the 1944 Completion Unit Layout Maps for Camp Beale, all three tanks were 12,000 gallon gasoline tanks. According to Beale AFB personnel, all three tanks were removed prior to 1962. The excavation pit is currently utilized as a compost pile for landscape waste.

This site was originally investigated by Law Environmental (LAW) in December 1994. LAW advanced seven (7) vapor points around the tank site and sampled the soil gas vapors. Analytical results of these samples established the presence of total petroleum hydrocarbons at concentrations as high as 27,000 parts per million by volume (ppmv) and benzene as high as 88 ppmv. Water was encountered in the tank pit area at six feet below ground surface. A visual inspection of the groundwater indicated the presence of free product. Soil gas sampling locations are shown on Figure 1. Analytical results are summarized in Table 1.

Table 1. Analytical Summary - Soil Gas Samples

Sample ID	Sample Depth (ft.bgs)	TPH Concentration (ppmv)	Benzene (ppmv)
SVP-132	6.5	27,120	6
SVP-133	7.5	226	1
SVP-134	8.5	ND	ND
SVP-142	8	23,620	88
SVP-143	7.5	12,030	15
SVP-144	7.5	1065	12
SVP-145	5.5	1	0.02

ppmv: Parts per million by volume ft.bgs: Feet below ground surface

As part of Metcalf and Eddy's investigation program, a total of seven (7) soil borings were advanced in the site vicinity to depths ranging from 6 feet to 18 feet below ground surface (bgs). Two of the borings were converted to monitoring wells (MW-1, MW-2). In addition, a total of seven (7) test pits (TP-8 through TP-14) were excavated to assist in the characterization process. The test pits were excavated to depths ranging from 6 feet to 9 feet bgs. Soil boring and test pit locations are shown on Figure 1.

During the field investigation at this site, groundwater was encountered in soil borings and test pits at depths ranging from 3 feet to 6 feet bgs. Photoionization detector (PID) readings observed ranged from <1 part per million (ppm) up to 2,500 ppm. The highest PID readings were observed n SB3, SB4, SB7, TP9, TP12, TP-13 and TP-14. Selected soil and groundwater samples were also collected and submitted to a state certified laboratory for analysis. The laboratory report is attached and summarized in Table 2.

Table 2. Analytical Summary - Metcalf & Eddy Site 10-494/10-495/10-496

				SOIL				
Sample ID			TPH-G (ug/g)	TPH-D (ug/g)	Benzene (ug/g)	Ethylbenzene (ug/g)	Xylenes (ug/g)	Total Lead (mg/kg)
SB1	02/21/95	6	<1	<1	< 0.005	< 0.005	< 0.005	1.2
SB1	02/21/95	16	<1	<1	<0.005	<0.005	< 0.005	2.9
SB2	02/22/95	18	<1	<1	< 0.005	<0.005	< 0.005	0.94
SB2	02/22/95	6.5	<1	<1	< 0.005	< 0.005	< 0.005	1.9
SB3	02/22/95	6	27	18	< 0.02	< 0.02	< 0.02	5.2
SB4	02/22/95	5.5	180	19	< 0.40	< 0.40	0.83	1.9
SB5	· 02/23/95	9	<1	<1	< 0.005	< 0.005	< 0.005	NA
SB5	02/23/95	6.5	<1	<1	< 0.005	<0.005	< 0.005	1.6
SB5	02/23/95	11	<1	<1	< 0.005	<0.005	< 0.005	1.5
SB6	02/23/95	5.5	<1	25	< 0.005	<0.005	< 0.005	4.8
SB7	02/23/95	5.5	<1	<1	< 0.005	<0.005	< 0.005	1.8
SB7	02/23/95	11.5	<1	<1	< 0.005	<0.005	< 0.005	1.4
Test Pit 10	03/29/95	6	<1	<1	< 0.005	< 0.005	< 0.005	1.6
Test Pit 11	Stained soil obs	erved at six	x feet bgs.	PID readings	s were > 100	0 ppm.		
Test Pit 12	Stained soil obs	erved at si	x feet bgs.	PID readings	s were > 100	0 ppm.		

	WATER									
Sample ID	Sample Date	Sample Depth (ft.bgs)	TPH-G (ug/L)	TPH-D (ug/L)	Benzene (ug/L)	Toluene (ug/L)	Ethyl- benzene (ug/L)	Xylenes (ug/L)	Total Lead (ug/L)	
MW-1	03/15/95	3.7	< 50	<50	<0.3	< 0.3	<0.5	<0.5	< 5.0	
SB3	02/22/95	5	5100	5800	4	10	8.6	50	NA	
Test Pit 8	03/29/95	6	290	140000	<0.6	<0.6	<1.0	<1.0	< 5.0	
Test Pit 9	03/29/95	6	4800	1600	3.8	2.4	4.9	11	< 5.0	
Test Pit 11	03/29/95	5.5	<50	< 50	<0.3	<0.3	< 0.5	<0.5	< 5.0	
Test Pit 12	03/29/95	8	370	830	<0.6	<1.0	<1.0	<1.0	7.4	
MW2	05/02/95	6	< 50	< 50	<0.3	<0.3	<0.3	<0.3	< 0.02	

ug/g: Micrograms per gram (parts per million)
ug/L: Micrograms per liter (parts per billion)

mg/kgs: Milligrams per kilogram (parts per million)

NA: Not analyzed

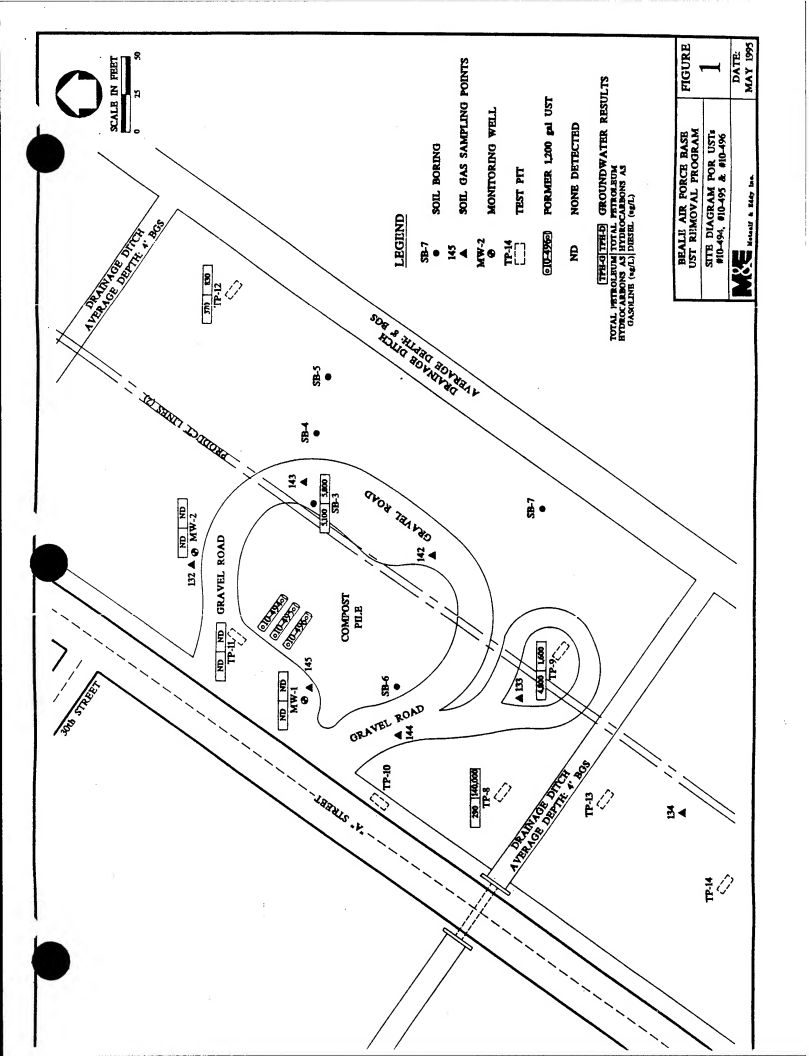
<: Less than (method detection limit)

>: Greater than

ft.bgs: Feet below ground surface

Based on the distribution of contaminants observed at this site, the westerly extent of contamination appears to be defined by TP-10, MW-1, and MW-2 (Figure 1). All other contaminant boundaries are presently undefined. Given the visual observations and PID readings observed in Test Pits 13 and 14, the southern extent of contamination appears to be extensive. The PID readings also indicate that the contamination has migrated either through or beneath the drainage ditch located south of the site. This ditch, as well as the other two ditches to the north and east, may affect the shallow groundwater flow direction in the immediate vicinity and provide a mechanism for contaminant migration.

The source of contamination at this site may be associated with the former tanks or the product line located east of the tanks which historically supplied fuel to several service stations along A street. Additional investigations at this site will be necessary to determine the source, define the full extent of contamination, and establish a groundwater flow direction and gradient.



LAW COMPANIES
Project # 221640003
BEALE AFB - Marysville, CA

ID:916-649-8100

TEG PROJECT #940822C

BROMODICHLOROMETHANE

cis-1,3 DICHLOROPROPENE

1,1,2 TRICHLOROETHANE

TETRACHLOROETHENE

BENZENE

TOLUENE

TPH

ETHYLBENZENE

TOTAL XYLENES

trans-1,3 DICHLOROPROPENE (ppmV)

Agreed Sample Love (appears to boxe free Product.

SAMPLE NUMBER	A001L 127VP	A001L 128VP	A001L 129VP	A001L 130VP	A001L) 132VP	A001L 133VP	A001L 134VP	A001L 135VP		
	0 01SV	001SV	001SV	001SV	001SV	0 015V	0015V	001SV		
COLLECTION DATE	9/26/94	9/26/94	9/26/94	9/28/94	9/26/94	8/26/94	9/25/94	9/25/94		
COLLECTION TIME	:	10:25	10:29	10:42	11:35	11:56	12:40	13:13	14:43	
COLLECTION DEPTH (feet)	13.5	7.5	7.5	13.5	6.5	8.0	8.5	7.5	_	
VINYL CHLORIDE	(ppmV)	nd	nd	nd	nd	nd	nd	nd	nd	
1,1 DICHLOROETHENE	(ppmV)	nd	nd	nd	nd	nd	nd	nd	nd	
METHYLENE CHLORIDE	(ppmV)	nd	nd	nd	nd	nd	nd	nd	กฮ	
trans-1,2 DICHLOROETHENE	(ppmV)	nd	nd	nd	nd	nd	nd	nd	nd	
1,1 DICHLOROETHANE	(ppmV)	nd	nd '	nd	nd	nd	nd	nd	nd	
cis-1,2 DICHLOROETHENE	(ppmV)	nd	nd	nd	nd	nd	nd	nd	nd	
CHLOROFORM	(ppniV)	nd	nd	nd	nd	nd	nd	nd	nd	
1,1,1 TRICHLOROETHANE	(ppmV)	nd	nd	nd	nd	nd	nd	nd	nd	
CARBON TETRACHLORIDE	(ppniV)	nd	nd	nd	nd	nd	nd	nd	nd	
2 DICHLOROETHANE	(ppmV)	nd	nd	nd	nd	nd	nd	nd	nd	
TRICHLOROETHENE	(ppmV)	nd	nd	nd	nd	nd	nd	nd	nd	
1,2 DICHLOROPROPANE	(ppmV)	nd	nd	nd	nd	nd	nd	nd	nd	

nd

EPA METHODS 8010, 8020 (BTEX) & mod8015 (TPH) ANALYSES OF SOIL XAPORS in ppmV

REPORTING LIMITS FOR ABOVE COMPOUNDS = 0.01 Part per Million by Volume (ppmV) (TPH & Vinyl Chloride = 'nd' NOT DETECTED AT LISTED REPORTING LIMITS

ANALYSES PERFORMED IN TEG'S DHS CERTIFIED MOBILE LAB (#1671)

(ppmV)

(ppmV)

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ANALYSES PERFORMED BY: Mr. Leif Jonsson & Ms. Stacie Wissler

DATA REVIEWED BY: Mr. Mark Jerphak

page 22

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Transglobal Environmental Geochemistry

PO Box 162580, Sacramento, CA 95816

Phone: (916) 736-3233

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Fax: (916) 452-5806



LAW COMPANIES Project # 221640003 BEALE AFB - Marysville, CA

TEG PROJECT #940822C

PA METHODS 8010, 8020 SAMPLE NUMBER:		A001L 136VP 001SV	A001L 137VP 001SV	A001L 13BVP 001SV	A001L 139VP 001SV	A001L 140VP 001SV	A001L 141VP 001SV	A001L 141VP 001SV dup	A001L 142VP 001SV
COLLECTION DATE: COLLECTION TIME: COLLECTION DEPTH (feet)		9/26/94 14:48 13.5	9/26/94 15:06 13.5	9/27/94 8:30 8.5	9/27/94 8:25 13.0	9/27/94 9;20 6.5	9/27/94 9:01 7.5	9/27/94 9:01 7.5	9/27/94 10:36 8 0
VINYL CHLORIDE 1,1 DICHLOROETHENE METHYLENE CHLORIDE trens-1,2 DICHLOROETHENE 1,1 DICHLOROETHANE	(ppmV) (ppmV) (ppmV)	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd
	(ppmV) (ppmV) (ppmV)	nd nd nd	nd nd nd	nd nd nd	nd nd nd -	nd nd nd	nd nd nd	nd nd nd	nd nd nd
	(ppmV) (ppmV)	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd
	(ppmV) (ppmV) (ppmV)	nd nd	nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd nd
	(ppmV) (ppmV) (ppmV)	nd nd nd	nd nd nd	nd nd	nd nd	nd nd nd	nd nd nd	nd nd nd	na na na
	(ppmV) -(ppmV) (ppmV)	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd 0.02	nd 0.24	nd 0.25	no
BENZENE TOLUENE	(ppmV) (ppmV)	nd nd	nd nd	nd nd	nd nd	nd nd	nd nd	nd nd nd	87. 36. 2.1
ETHYLBENZENE TOTAL XYLENES	(ppmV) (ppmV) (ppmV)	nd nd nd	nd nd nd	nd nd nd	nd nd nd	nd nd 3	nd nd nd	na nd nd	5.6 236

REPORTING LIMITS FOR ABOVE COMPOUNDS = 0.01 Part per Million by Volume (ppmV) (TPH & Vinyl Chloride = 1 ppmV)

'nd' NOT DETECTED AT LISTED REPORTING LIMITS

ANALYSES PERFORMED IN TEG'S DHS CERTIFIED MOBILE LAB (#1671)

ANALYSES PERFORMED BY: Mr. Leif Jonsson & Ms. Stacie Wissler

DATA REVIEWED BY: Mr. Mark Jerphak

page 23

Transglobal Environmental Geochemistry

PO Box 162580, Sacramento, CA 95816

Phone: (916) 736-3233

Fax: (916) 452-5806



LAW COMPANIES Project # 221640003 BEALE AFB - Marysville, CA

TEG PROJECT #940822C

SAMPLE NUMBER: COLLECTION DATE: COLLECTION TIME:		A001L 143VP 001SV 8/27/94 10:36	A001L 144VP 001SV 9/27/94 10:53	A001L 145VP 001SV 9/27/94 11:02 7.0	A001L 145VP 002SV 9/27/94 11:25 5.5	A001L 146VP 001SV 9/27/94 14:13 8.5	A001L 147VP 001SV 8/27/94 14:45 10.5	A001L 147VP 001SV dup 9/27/94 14:45 10.5	A001L 147VP 003SV 9/27/94 15:07 5.0						
										COLLECTION DEPTH (feet)	COLLECTION DEPTH (feet):		7.5		
										VINYL CHLORIDE	(ppmV)	nd	nd	nd	nd
1,1 DICHLOROETHENE	(ppmV)	nd	nd							nd	nd	nd		nd nd	nd
METHYLENE CHLORIDE	(ppmV)	nd	nd	nd	nd	nd	nd nd	nd and	nd						
trans-1,2 DICHLOROETHENE	(ppmV)	nd	nd	nd	nd	nd	nd ===	nd 5.d	nd						
1,1 DICHLOROETHANE	(ppmV)	nd	nd	nd	nd		лd ~-d	nd 	nd						
cis-1,2 DICHLOROETHENE	(ppmV)	nd	nd	nd	nd nd	nd nd	nd nd	nd	nd 						
CHLOROFORM	(ppmV)	nd	nd	nd	nd nd	na nd	nd = d	nd	nd						
1.1,1 TRICHLOROETHANE	(ppmV)	nd	nd	nd	nd		nd nd	nd 	nd						
CARBON TETRACHLORIDE	(ppniV)	nd	nd	nd nd	na nd	nd nd	nd nd	nd ma	nd						
,2 DICHLOROETHANE	(ppmV)	nd	nd	nd	nd		nd ad	nd 	nd						
RICHLOROETHENE	(ppmV)	nd	nd	nd nd	na nd	nd nd	nd 0.08	nd 0 .07	nd						
1.2 DICHLOROPROPANE	(ppmV)	nd	nd	nd	nd				nd						
ROMODICHLOROMETHANE	(ppmV)	nd	nd	nd nd	· · ·	nd nd	nd	nd	nd						
is-1,3 DICHLOROPROPENE	(ppmV)	กซ	nd	na nd	กส	nd	nd	nd	nd						
rans-1,3 DICHLOROPROPENE	(ppmV)	nd nd	nd nd		nd	nd 	nd	nd	nd						
1,1,2 TRICHLOROETHANE	(ppmV)	nd nd	nd nd	กช กช	กฮ	nd a.d	nd	nd	nd						
TETRACHLOROETHENE	(ppmV)	nd	nd	nd nd	nd ad	nd	nd O.11	nd 0.10	na						
	(PP/// V)	710		nd 	nd	nd	0.11	0.12	nd						
ENZENE	(ppmV)	15.35	11.72	0.02	nd	nd	nd	nd	nd						
OLUENE	(ppmV)	3.84	4.17	0.04	nd	0.11	nd	nd	nd						
THYLBENZENE	(ppmV)	0.09	0.35	0.06	nd	0.02	nd	nd	nd						
TOTAL XYLENES	(ppmV)	0.57	1.92	0.13	nd	0.11	nd	nd	nd						
TPH	(ppmV)	12030	1065	7	1	10	nd	nd	nd nd						

REPORTING LIMITS FOR ABOVE COMPOUNDS = 0.01 Part per Million by Volume (ppmV) (TPH & Vinyl Chloride = 1 ppmV) 'nd' NOT DETECTED AT LISTED REPORTING LIMITS

ANALYSES PERFORMED IN TEG'S DHS CERTIFIED MOBILE LAB (#1671)

ANALYSES PERFORMED BY: Mr. Lelf Jonsson & Ms. Stacie Wissler

DATA REVIEWED BY: Mr. Mark Jerpbak

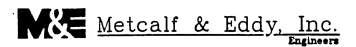
page 24

Transglobal Environmental Geochemistry

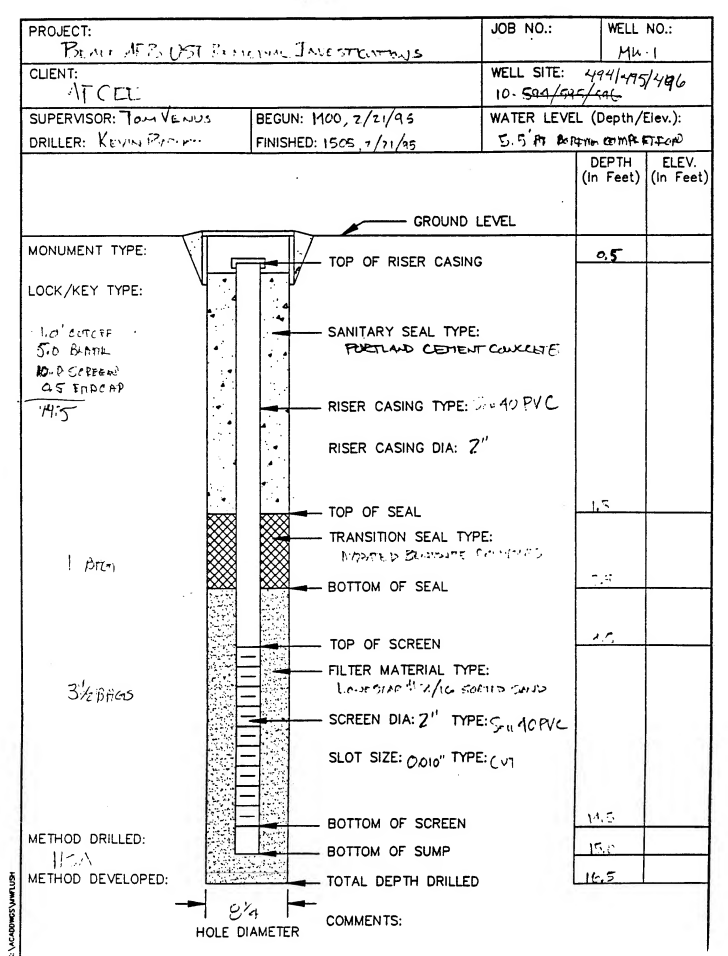
PO Box 162580, Sacramento, CA 95816

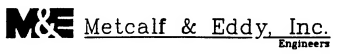
Phone: (916) 736-3233

Fax: (916) 452-5806



MONITORING WELL CONSTRUCTION DIAGRAM





GEOLOGIC LO

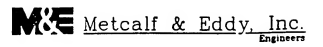
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DR	RILLIN			D: 	nt or		DRILL		FLUID JE	/S0U	RCE:	TOP OF ROCK	-	/ELEV.):
DEPTH	SAMPLE TYPE/NO.	SAMPLE DEPTH	SAMPLE RECOVERY	BLOW COUNT (per 6 inches)	PERCENT RECOVERY	PRODU			TIME LOG	USCS LOG		STRATIC DESCR	GRAPHIC SIPTION	
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	DEPTH	SAMPLE TYPE/NO.	SAMPLE DEPTH	SAMPLE RECOVERY	BLOW COUNT per 6 inches)	PID READING	PRODU OVA MEA	ASURM TC.	ENT,	=	NSCS LOG			STRAT	RIPTIO	NC	7	
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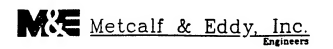
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GEOLOGIC BORING LOG

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GEOLOGIC BORING LOG

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APPENDIX C

LABORATORY ANALYTICAL DATA



Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 94710, Phone (510) 486-0900

ANALYTICAL REPORT

Prepared for:

Parsons Engineering Science, Inc. 1301 Marina Village Parkway Suite 200 Alameda, CA 94501

Date: 18-MAR-96

Lab Job Number: 124633

Project ID: 722450.14020 Location: Beale AFB

Reviewed by: Tursa k Morriso

Reviewed by:

This package may be reproduced only in its entirety.



Laboratory Number: 124633 sa

Client: Parsons Engineering Science

Project#: 722450.14020 Location: Beale AFB Sample Date: 02/27/96 Receipt Date: 02/28/96

CASE NARRATIVE

Curtis & Tompkins Ltd. received five soil samples from the above referenced project on Feb.28,1996. All samples were received cold and intact. Chromatograms are included in this report for any samples containing quantifiable concentrations of hydrocarbons, along with an example chromatogram for each type of fuel standard. The following analytical problems were encountered:

TPH/D: No analytical problems.

TPH/G: Sample chromatograms do not resemble Curtis & Tompkins' gasoline standard, but are similar to a JP-4 standard which is occasionally requested.

BTXE: The qualifier "C" flag appears with results for samples BE-MP14-7 (C&T# 124633-004) and BE-MP22-7 (C&T# 124633-005). This flag indicates that the target compound was confirmed by a second analytical column, but that the quantitated result on the confirmation column differed from the reported result by more than a factor of two. This often happens when the observed hydrocarbon pattern does not resemble gasoline, which is the case with this sample set. The matrix spike and spike duplicate failed ethylbenzene and m,p-xylene recovery limits. The spiked sample was not from Beale AFB. The associated laboratory control sample passed all acceptance criteria.



TEH-Tot Ext Hydrocarbons

client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: LUFT

Location: Beale AFB

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124633-001 BE-MP09-6	26268	02/27/96	03/05/96	03/09/96	14%
124633-002 BE-MP12-4	26268	02/27/96	03/05/96	03/09/96	16%
124633-003 BE-MP13-6.5	26268	02/27/96	03/05/96	03/09/96	14%
124633-004 BE-MP14-7	26268	02/27/96	03/05/96	03/09/96	15%

Analyte Diln Fac:	Units	124633-001 1	124633-002 1	124633-003 1	124633-004 1
JP5	mg/Kg	2.3	<1.2	7 Y	55 Y
Diesel Range	mg/Kg	8.8YH	<1.2	4.7YL	33 YL
Motor Oil Range	mg/Kg	15 YL	<6	<5.8	21 YL
Surrogate					
exacosane	%REC	99	93	92	94

Y: Sample exhibits fuel pattern which does not resemble standard

H: Heavier hydrocarbons than indicated standard

L: Lighter hydrocarbons than indicated standard



TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: LUFT

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124633-005 BE-MP22-7	26268	02/27/96	03/05/96	03/09/96	20%

Analyte Diln Fac:	Units	124633- 1	-005			
JP5	mg/Kg	340	YH	 		
Diesel Range	mg/Kg		YL			
Motor Oil Range	mg/Kg	23	YL			
Surrogate						
Hexacosane	%REC	84		 		

- Y: Sample exhibits fuel pattern which does not resemble standard
- H: Heavier hydrocarbons than indicated standard
- L: Lighter hydrocarbons than indicated standard



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Parsons Engineering Science, Inc. lient:

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method:

LUFT

Location: Beale AFB

Soil

METHOD BLANK

Prep Date:

03/05/96

Batch#: 26268 Units: mg/Kg Analysis Date:

03/08/96

Diln Fac: 1

Matrix:

MB Lab ID: QC16519

Result	
<1.0	
<1.0	
<5.0	
%Rec	Recovery Limits
69	60-140
	<1.0 <1.0 <5.0



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: LUFT

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Matrix: Soil

Prep Date:

03/05/96

Batch#: 26268 Units: mg/Kg Analysis Date: 03/08/96

Diln Fac: 1

LCS Lab ID: QC16520

Analyte	Result	Spike Added	%Rec #	Limits
Diesel Range	40.1	49.5	81	60-140
Surrogate	%Rec	Limits		
Hexacosane	69	60-140		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: LUFT

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Lab ID: 124641-004

02/27/96 Sample Date: Received Date: 03/01/96

Matrix: Soil

Prep Date: 03/05/96 Analysis Date:

Batch#: 26268

03/08/96

Units: mg/Kg dry weight

Diln Fac: 1

Moisture:

35%

MS Lab ID: QC16521

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Diesel Range	76.15	<1.538	71.69	94	60-140
Surrogate	%Rec	Limits			
Hexacosane	82	60-140			

MSD Lab ID: QC16522

alyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Diesel Range	76.15	63.54	83	60-140	12	<30
Surrogate	%Rec	Limit	s		<u> </u>	•
Hexacosane	74	60-14	0			

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

GC15 Channel A Surrogate

Sample Name : S,124633-001,26268
FileName : C:\GC15\CHB\068B015.raw

: DUAL Method

Start Time : 0.00 min

End Time : 31.90 min Plot Offset: 32 mV

Sample #: 26268

Date: 3/9/96 03:48 AM

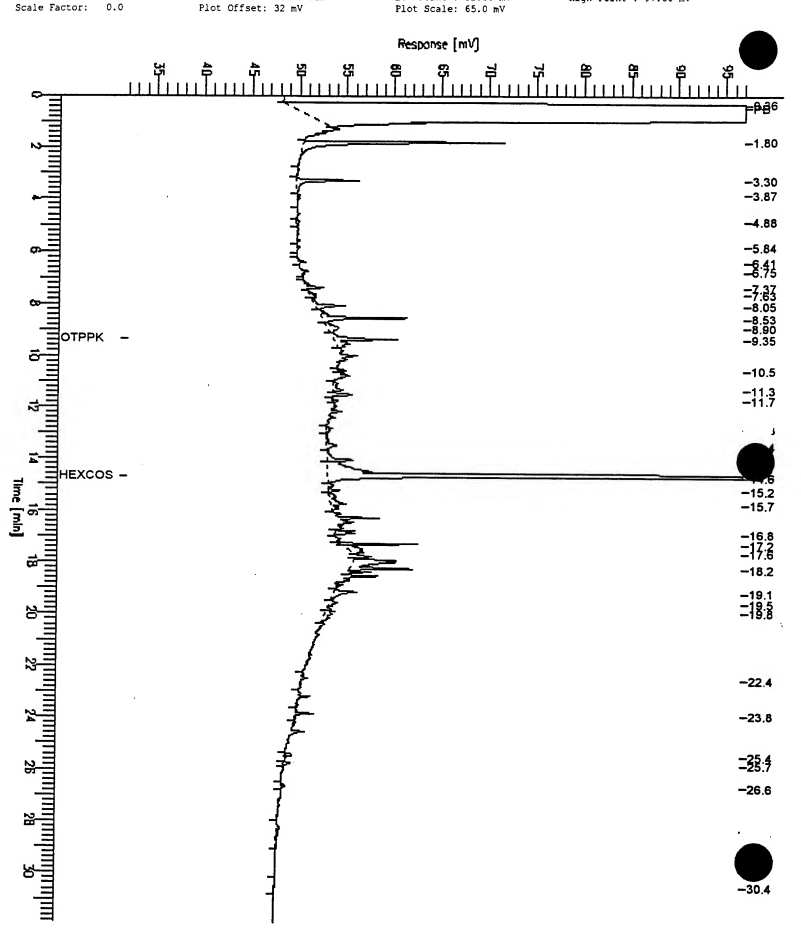
Time of Injection: 3/9/96 03:15 AM

Low Point : 32.00 mV

High Point : 97.00 mV

Page 1 of 1

Plot Scale: 65.0 mV



GC15 Channel A Surrogate

Sample Name: S,124633-003,26268

: C:\GC15\CHB\068B012.raw FileName

Method : DUAL Start Time : 0.00 min

e Factor: 0.0 End Time : 31.90 min Plot Offset: 32 mV

Page 1 of 1

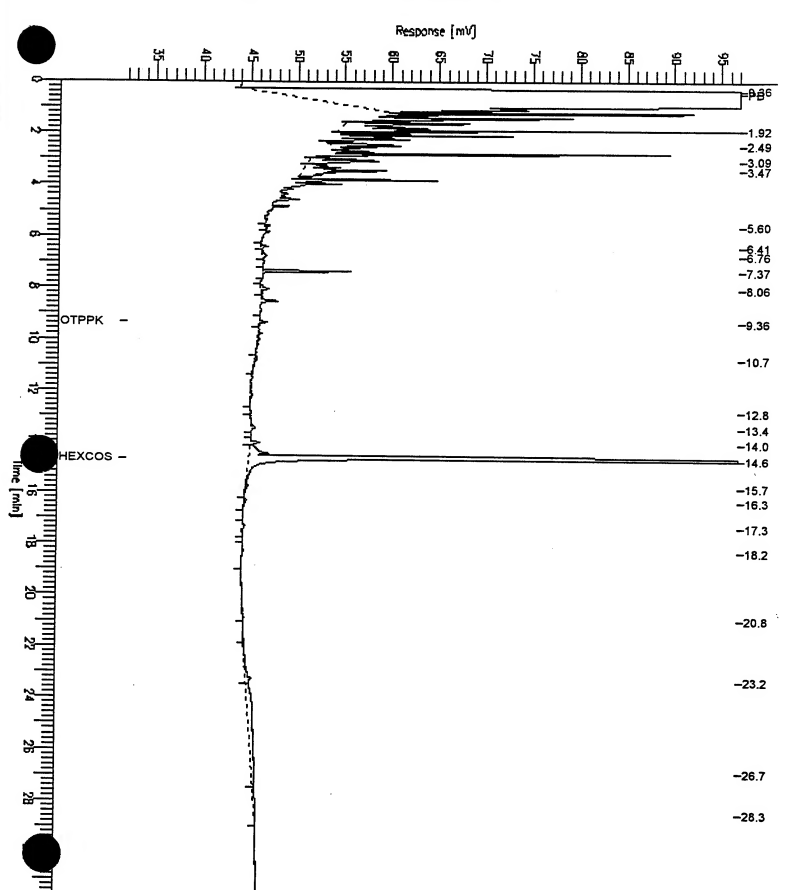
Sample #: 26268 Date: 3/9/96 01:37 AM

Time of Injection: 3/9/96 01:04 AM

Low Point : 32.00 mV

High Point: 97.00 mV

Plot Scale: 65.0 mV



GC15 Channel A TEH

Sample Name: S,124633-004,26268 FileName : C:\GC15\CHB\068B021.RAW

Method : BTEH.MTH

Start Time : 0.01 min Scale Factor: 0.0

End Time : 31.91 min Plot Offset: 43 mV

Sample #: 26268

Date: 3/11/96 03:50 PM Time of Injection: 3/9/96

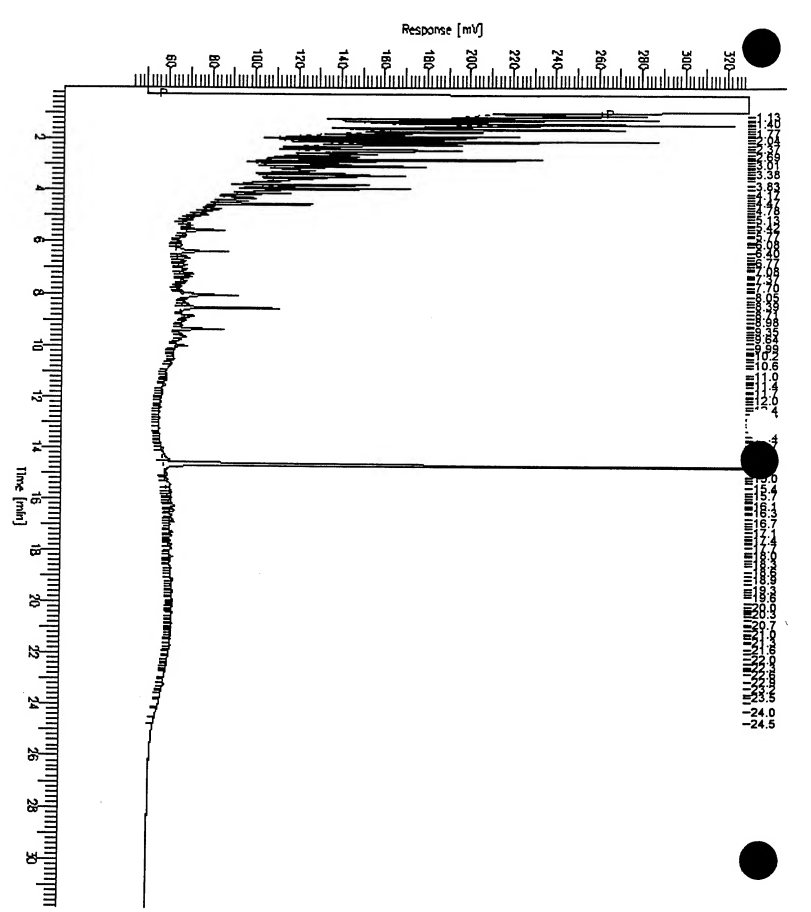
07:38 AM

High Point: 329.56 mV

Page 1 of 1



Plot Scale: 286.5 mV



GC15 Channel A TEH

Sample Name : S,124633-005,26268

FileName : C:\GC15\CHB\071B005.RAW

Method : BTEH.MTH Start Time : 0.01 min

End Time

Factor: 0.0 : 31.91 min

Plot Offset: 30 mV

Sample #: 26268

Date: 3/11/96 06:32 PM

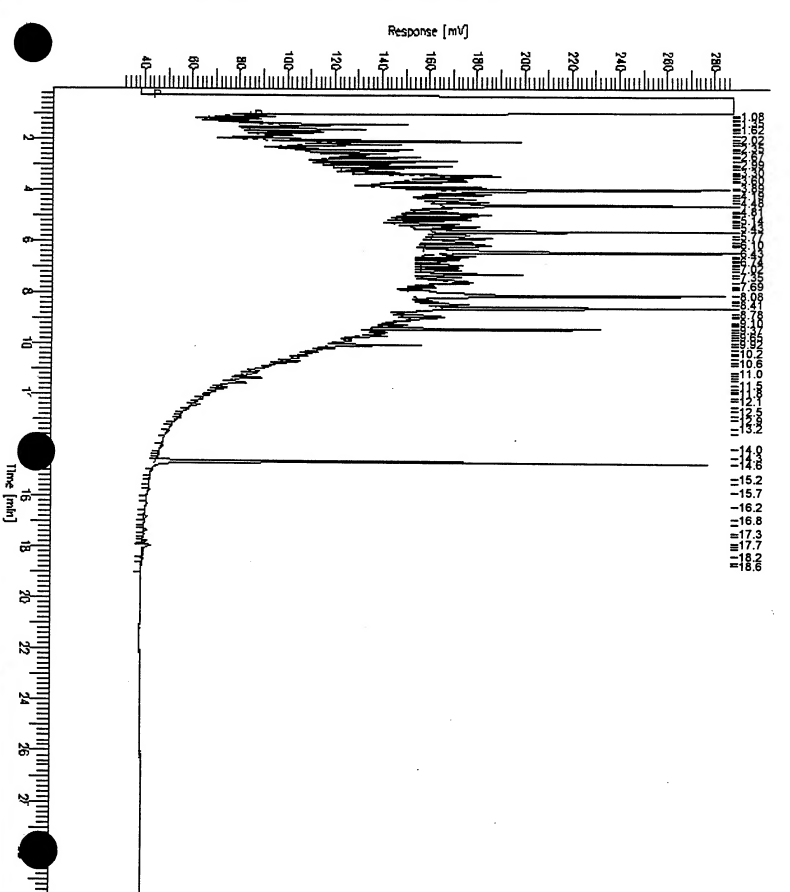
Time of Injection: 3/11/96 05:11 PM

Low Point: 30.24 mV

High Point : 287.89 mV

Page 1 of 1

Plot Scale: 257.7 mV



GC15 Channel A TEH

Sample Name : CCV, JP5
FileName : C:\GC15\CHB\067B023.RAW

: BTEH.MTH Method

Start Time : 0.01 min

End Time : 26.87 min

Scale Factor: 0.0

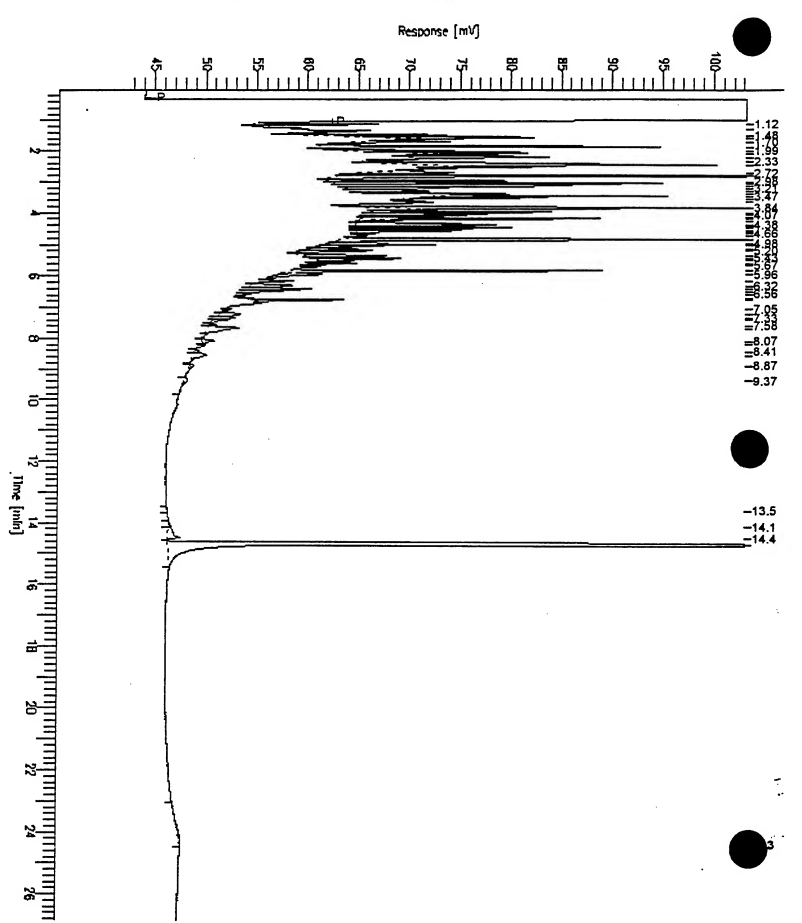
Plot Offset: 42 mV

Sample #: 250MG/L Date: 3/8/96 01:50 PM Page 1 of 1

Time of Injection: 3/8/96 07:20 AM

High Point : 103.22 mV

Low Point: 42.42 mV Plot Scale: 60.8 mV



DIESEL

GC15 Channel A TEH

Sample Name : CCV, 96WS1948, DSL

: C:\GC15\CHB\067B021.RAW FileName

: BTEH.MTH Method

- Time : 0.01 min

End Time : 31.91 min

Plot Offset: 42 mV

Sample #: DSL

Page 1 of 1

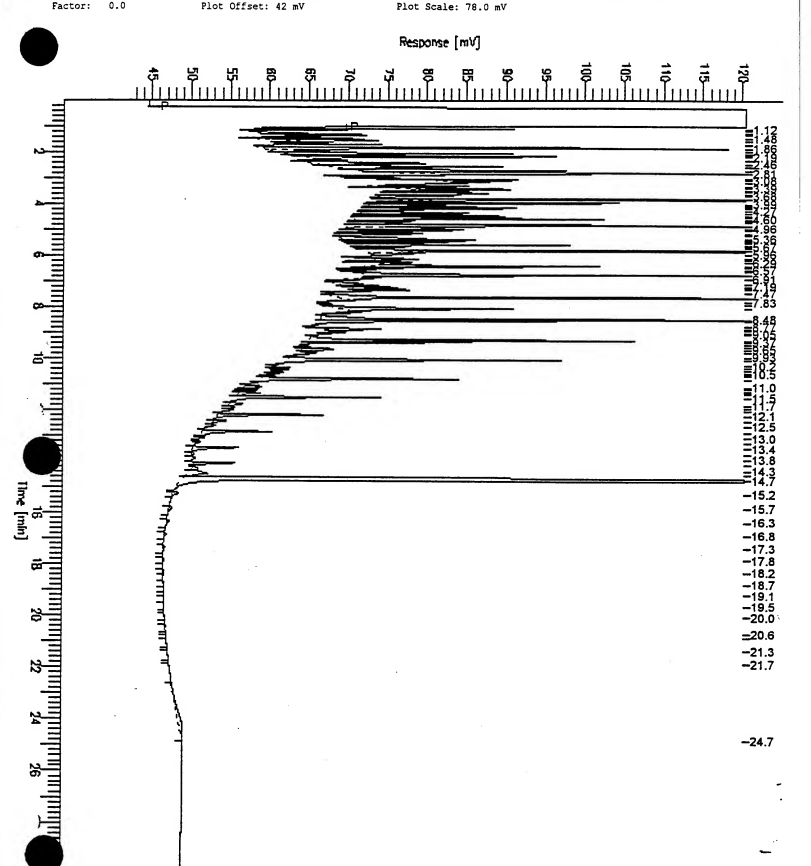
Date: 3/8/96 01:45 PM

Time of Injection: 3/8/96 05:52 AM

Low Point: 42.49 mV

Plot Scale: 78.0 mV

High Point : 120.49 mV



GC15 Channel A TEH

Sample Name: MO 95W5/596
FileName: C:\GC15\CHB\067B008.RAW
Method: BTEH.MTH
Start Time: 0.01 min

Scale Factor: 0.0

: 31.91 min Plot Offset: 42 mV

50ong)

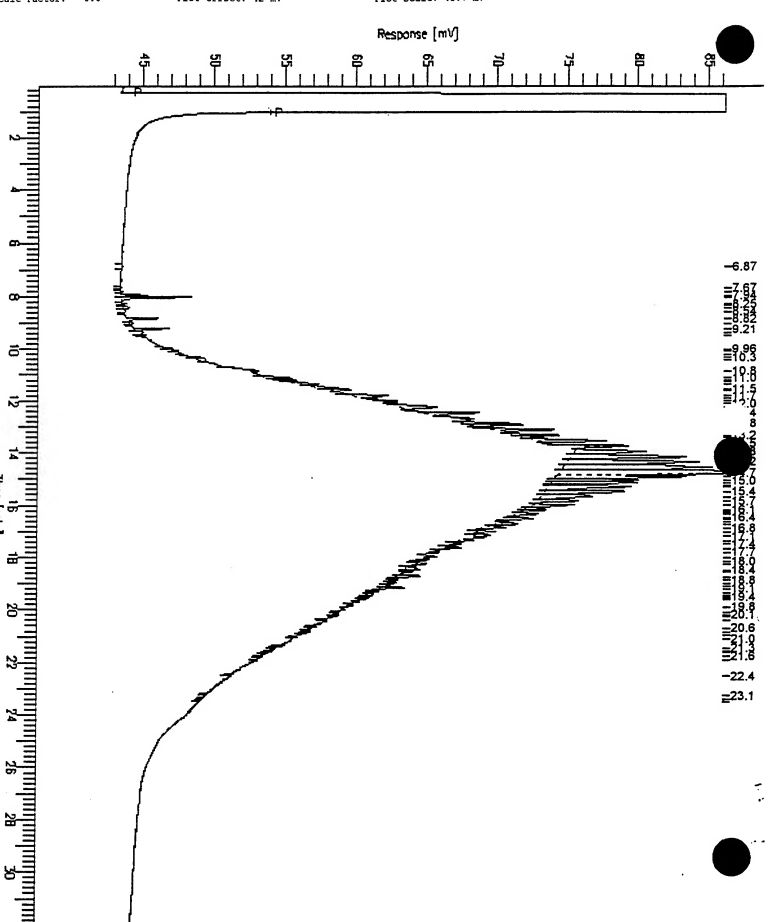
Sample #: Date: 3/8/96 02:01 PM Time of Injection: 3/7/96 08:23 PM

Low Point: 42.50 mV

High Point : 86.16 mV

Page 1 of 1

Plot Scale: 43.7 mV



TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

roject#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124633-001 BE-MP09-6	26345	02/27/96	03/12/96	03/12/96	14%
124633-002 BE-MP12-4	26345	02/27/96	03/12/96	03/12/96	16%
124633-003 BE-MP13-6.5	26345	02/27/96	03/12/96	03/12/96	14%
124633-004 BE-MP14-7	26345	02/27/96	03/12/96	03/12/96	15%

Analyte Diln Fac:	Units	124633-001 1	124633-002 1	124633-003 1	124633-004 1
Gasoline	mg/Kg	<1.2	<1.2	<1.2	21 Y
Surrogate					
Trifluorotoluene	%REC	89	94	67	115
Rromobenzene	%REC	79	84	58	73

Sample exhibits fuel pattern which does not resemble standard



TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124633-005	BE-MP22-7	26389	02/27/96	03/12/96	03/12/96	20%

Analyte Diln Fac:	Units	124633-0 25	05
Gasoline	mg/Kg	44	YH
Surrogate			
Trifluorotoluene	%REC	92	
Bromobenzene	%REC	65	

Y: Sample exhibits fuel pattern which does not resemble standard

H: Heavier hydrocarbons than indicated standard



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

client: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Method: EPA 5030

Location: Beale AFB

METHOD BLANK

Matrix: Soil Prep Date: 03/12/96

Batch#: 26389

Units: mg/Kg

Diln Fac: 1

Analysis Date: 03/12/96

MB Lab ID: QC16980

Analyte	Result	
Gasoline	<1.0	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene Bromobenzene	96 85	52-127 47-112



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015)

Project#: 722450.14020

Soil

Prep Method: EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Prep Date: 03/11/96 Analysis Date: 03/11/96

Batch#: 26345 Units: mg/Kg

Matrix:

Units: mg/I Diln Fac: 1

LCS Lab ID: QC16892

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	10.1	10	101	80-120
Surrogate	*Rec	Limits		
Trifluorotoluene Bromobenzene	106 90	52-127 47-112		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits



TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124634-001	BE-MP15-9.5	26389	02/27/96	03/12/96	03/12/96	10%
124634-002	BE-MP10-5.5	26389	02/28/96	03/12/96	03/12/96	10%
124634-003	BE-MP11-7.0	26389	02/28/96	03/12/96	03/12/96	12%
124634-004	BE-MP19-7.0	26345	02/28/96	03/12/96	03/12/96	9%

Analyte Diln Fac:	Units	124634-001 1	124634-002 200	124634-003 25	124634-004 1
Gasoline	mg/Kg	22 Y	780 YH	40 YH	<1.1
Surrogate					
rifluorotoluene	%REC	90	109	67	89
romobenzene	%REC	68	77	56	78

Y: Sample exhibits fuel pattern which does not resemble standard

H: Heavier hydrocarbons than indicated standard



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

METHOD BLANK

Matrix: Soil

Prep Date: 03/11/96 Analysis Date: 03/11/96

Batch#: 26345 Units: mg/Kg Diln Fac: 1

MB Lab ID: QC16891

Analyte	Result	
Gasoline	<1.0	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene	93	52-127
Bromobenzene	78	47-112

GC15 Channel A TEH

Sample Name : CCV, 96WS1948, DSL

: C:\GC15\CHB\067B021.RAW FileName

: BTEH.MTH Method

> ` Time : 0.01 min : Factor: 0.0

End Time : 31.91 min

Plot Offset: 42 mV

Sample #: DSL

Date: 3/8/96 01:45 PM

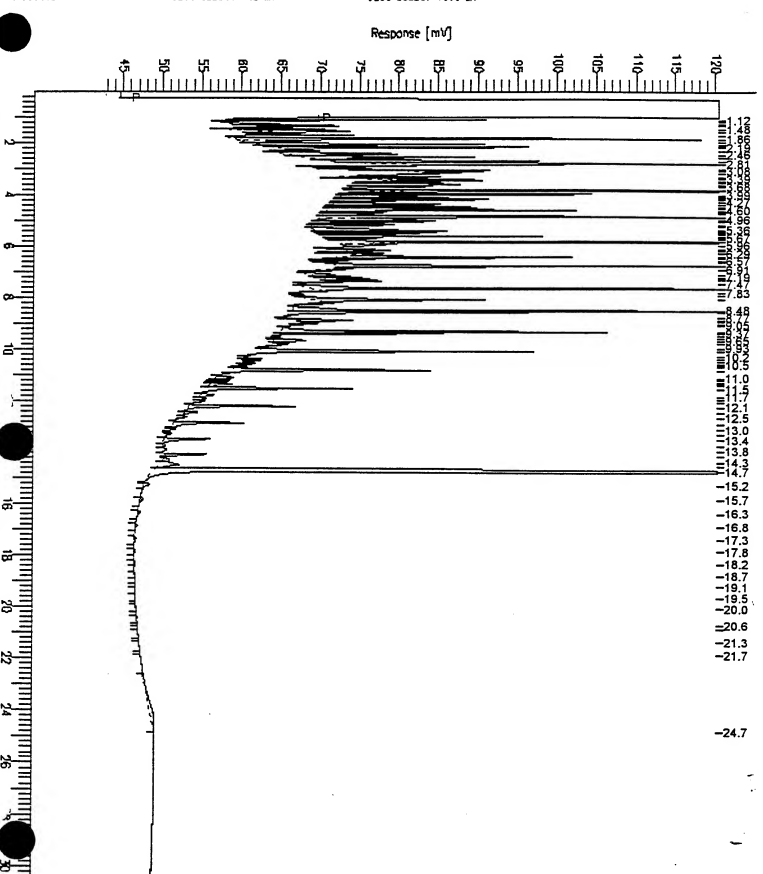
Time of Injection: 3/8/96 05:52 AM

Low Point: 42.49 mV

High Point: 120.49 mV

Page 1 of 1

Plot Scale: 78.0 mV



mo 95ws/596 Sample Name :

: C:\GC15\CHB\067B008.RAW : BTEH.MTH FileName

Method

Start Time : 0.01 min Scale Factor: 0.0

End Time : 31.91 min Plot Offset: 42 mV

Sample #: Date: 3/8/96 02:01 PM

Time of Injection: 3/7/96

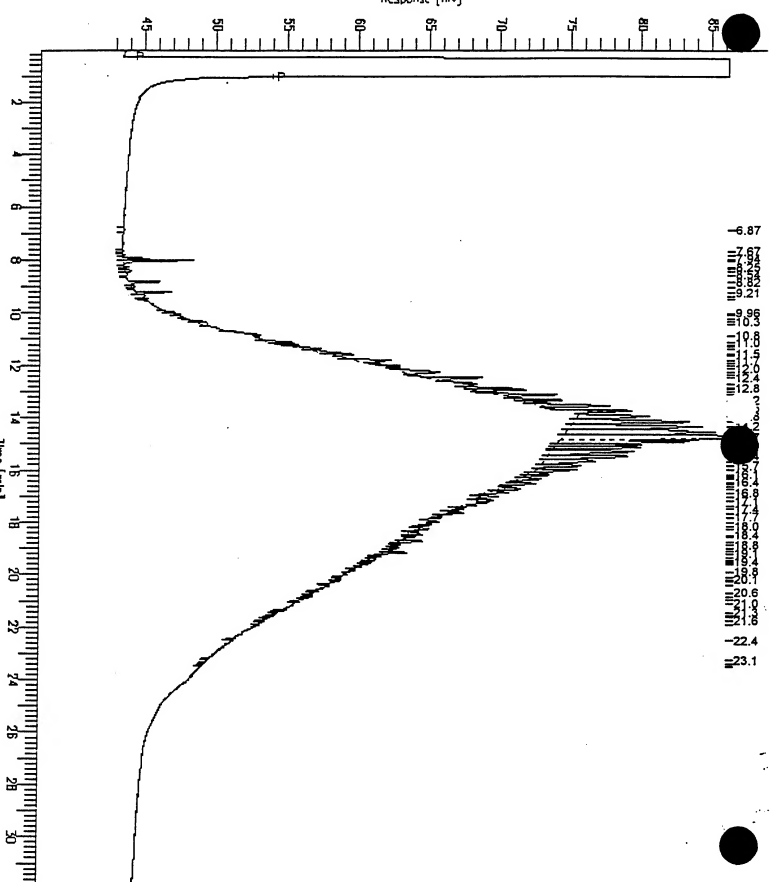
08:23 PM

High Point : 86.16 mV

Page 1 of 1

Low Point: 42.50 mV Plot Scale: 43.7 mV





GC15 Channel A Surrogate

Sample Name : S,124634-004,26268

FileName : C:\GC15\CHB\068B008.raw

: DUAL Method Start Time : 0.00 min

e Factor: 0.0

End Time : 31.90 min

Plot Offset: 32 mV

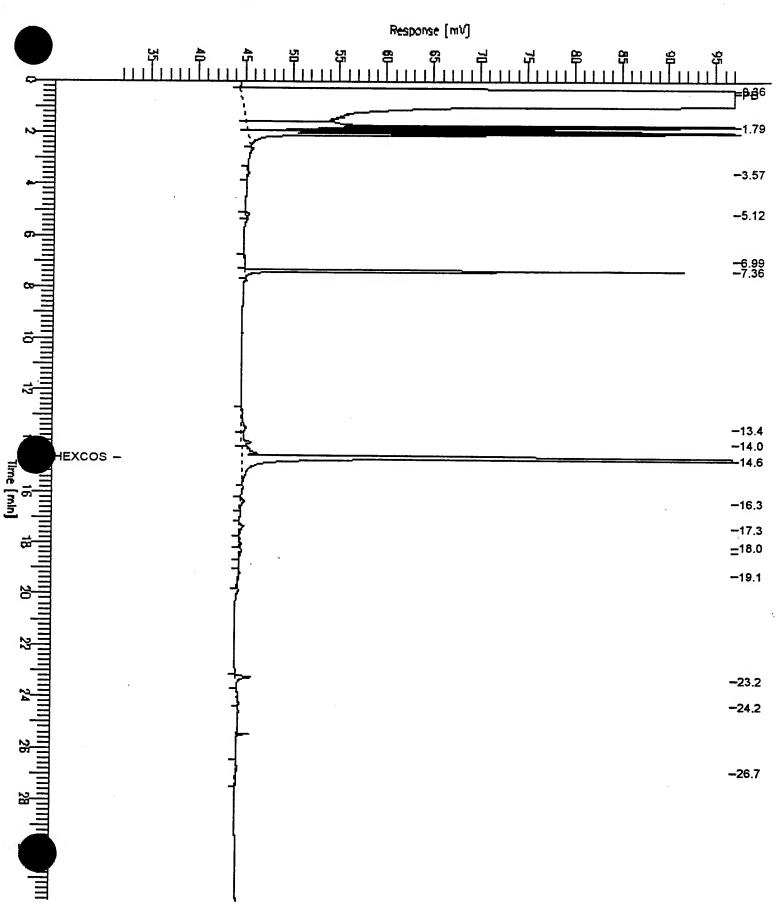
Sample #: 26268 Date: 3/8/96 10:42 PM

Time of Injection: 3/8/96 10:09 PM

High Point : 97.00 mV Low Point : 32.00 mV

Page 1 of 1

Plot Scale: 65.0 mV



JP5

GC15 Channel A TEH

Sample Name : CCV, JP5

: C:\GC15\CHB\067B023.RAW FileName

Method : BTEH.MTH

Start Time : 0.01 min Scale Factor: 0.0

End Time : 26.87 min

Plot Offset: 42 mV

Sample #: 250MG/L Date: 3/8/96 01:50 PM

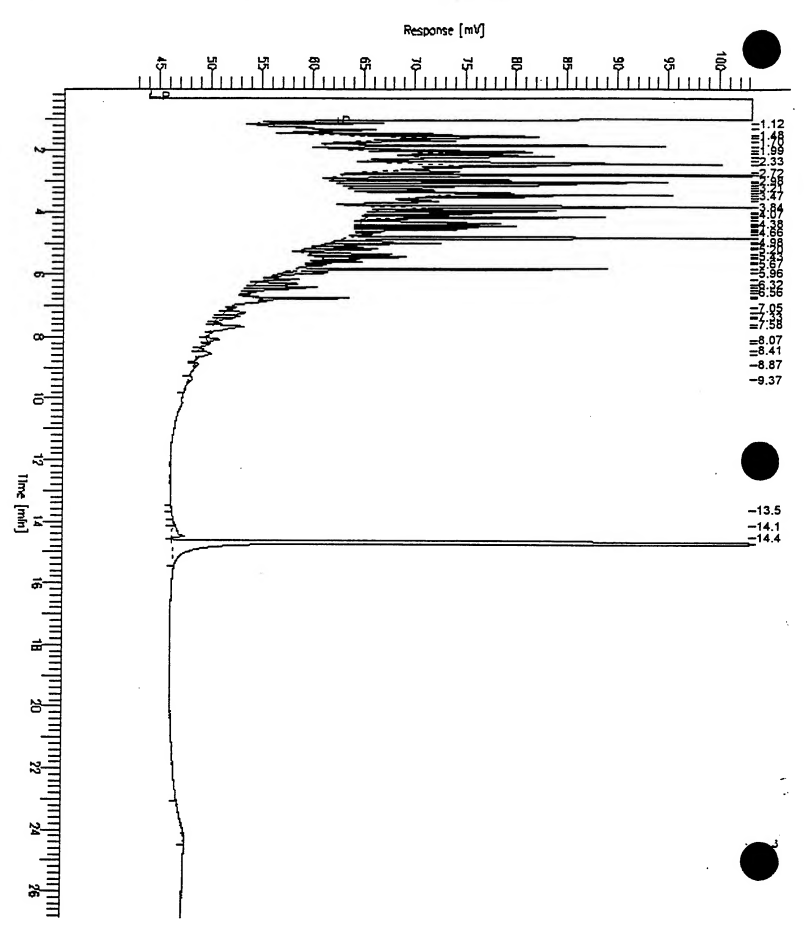
Time of Injection: 3/8/96 07:20 AM

Low Point : 42.42 mV

High Point : 103.22 mV

Page 1 of 1

Plot Scale: 60.8 mV



GC15 Channel A TEH

Sample Name: S,124634-002,26268

FileName : C:\GC15\CHB\068B007.RAW

Method : BTEH.MTH Start Time : 0.01 min

End Time : 31.91 min

· Factor: 0.0 Plot Offset: 38 mV

Sample #: 26268 Date: 3/11/96 02:54 PM

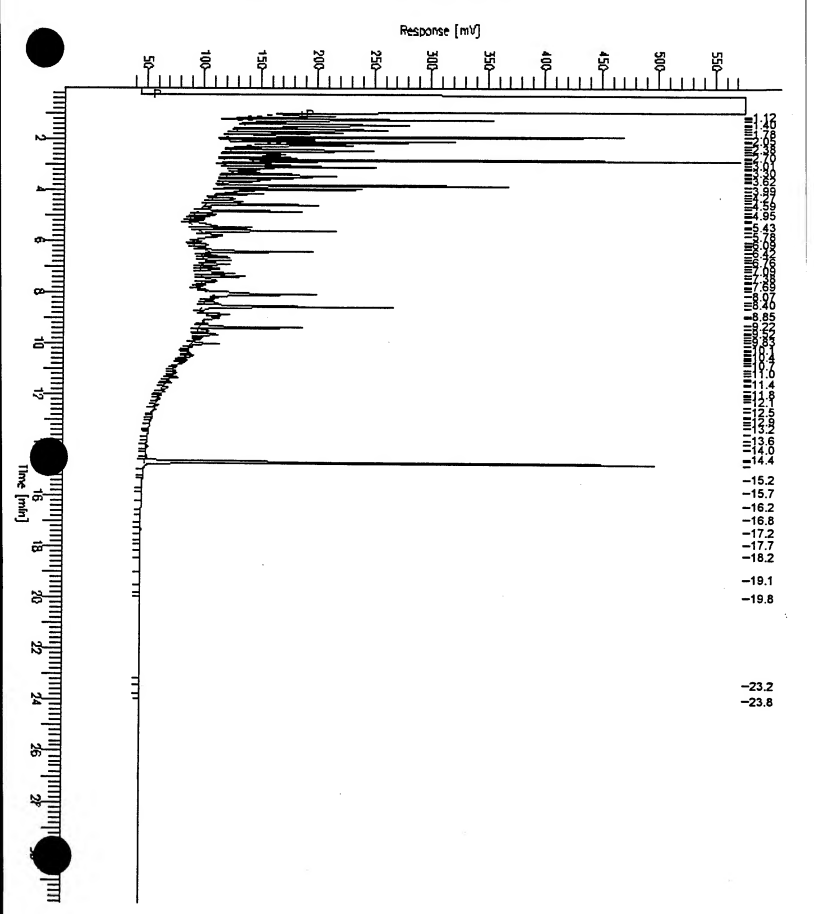
Time of Injection: 3/8/96 09:26 PM

Low Point: 37.87 mV

High Point : 576.33 mV

Page 1 of 1

Plot Scale: 538.5 mV



GC15 Channel A Surrogate

Sample Name: S,124634-003,26268

: C:\GC15\CHB\068B006.raw FileName

Method : DUAL

Start Time : 0.00 min

Scale Factor: 0.0 End Time : 31.90 min

Plot Offset: 32 mV

Sample #: 26268

Date: 3/8/96 09:14 PM

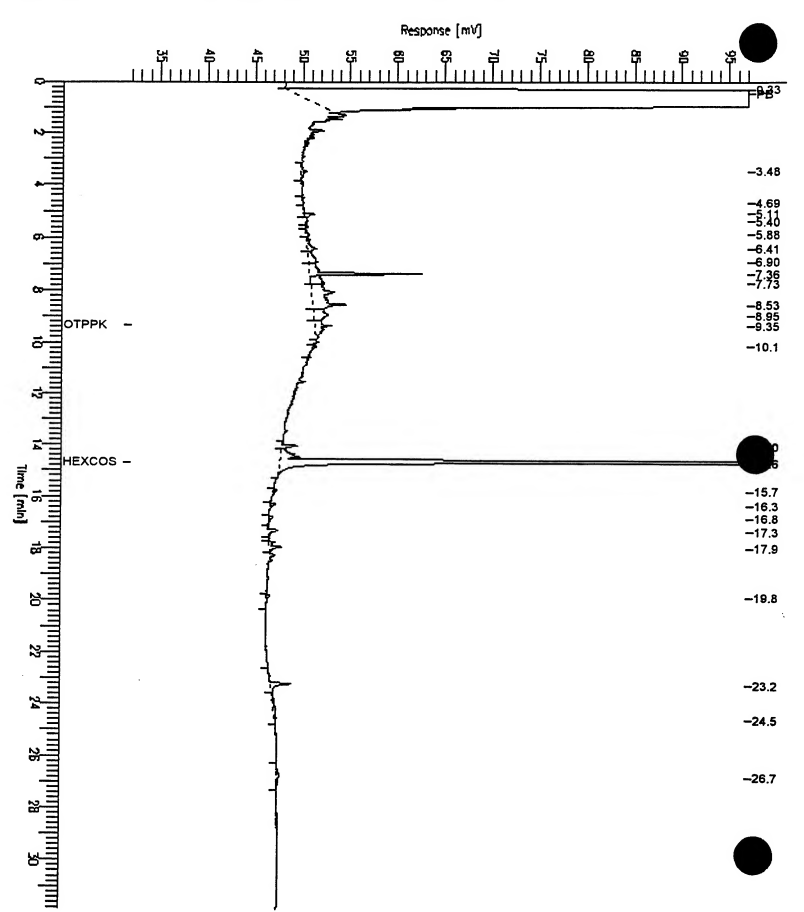
Time of Injection: 3/8/96 08:42 PM

Low Point : 32.00 mV

High Point : 97.00 mV

Page 1 of 1

Plot Scale: 65.0 mV



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

:lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Location: Beale AFB

Prep Method: LUFT

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ 124641-004

Sample Date: Received Date:

02/27/96 03/01/96

Lab ID: Matrix: Soil

Prep Date: Analysis Date: 03/05/96 03/08/96

Batch#: 26268

Units: mg/Kg dry weight

Moisture:

35%

Diln Fac: 1

MS Lab ID: QC16521

Analyte	Spike Added	Sample	MS	%Rec #	Limits	
Diesel Range	sel Range 76.15		71.69	94	60-140	
Surrogate	%Rec	Limits		<u></u>		
Hexacosane	82	82 60-140				

MSD Lab ID: QC16522

nalyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Diesel Range	76.15	63.54	83	60-140	12	<30
Surrogate	%Rec	Limits				
Hexacosane	74	60-140				

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

Sample Name : S,124634-001,26268 FileName : C:\GC15\CHB\068B014.RAW

Method : BTEH.MTH

Start Time : 0.01 min Scale Factor: 0.0

End Time

: 31.91 min Plot Offset: 42 mV

Sample #: 26268 Date: 3/11/96 02:53 PM

Time of Injection: 3/9/96

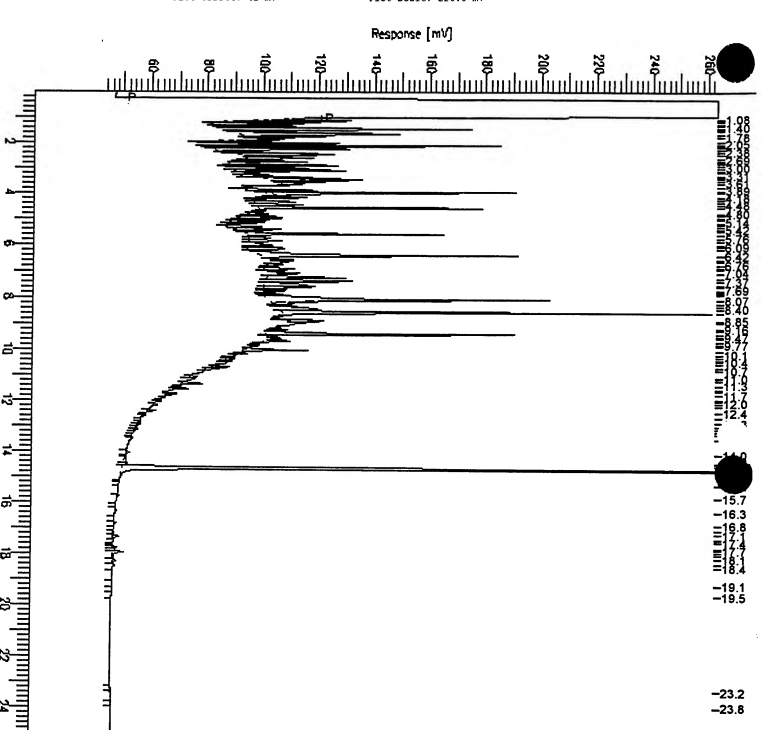
Low Point: 42.46 mV

02:32 AM

Plot Scale: 220.8 mV

High Point : 263.26 mV

Page 1 of 1





BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Parsons Engineering Science, Inc. Client:

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method:

Location: Beale AFB

LUFT

METHOD BLANK

Matrix: Soil Prep Date:

03/05/96

Batch#: 26268 Units: mg/Kg Analysis Date:

03/08/96

Diln Fac: 1

MB Lab ID: QC16519

Analyte	Result	
JP5	<1.0	
Diesel Range	<1.0	
Motor Oil Range	<5.0	
Surrogate	%Rec	Recovery Limits
Hexacosane	69	60-140



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method:

LUFT

LABORATORY CONTROL SAMPLE

Matrix: Soil Batch#: 26268 Units: mg/Kg

Prep Date: Analysis Date: 03/05/96 03/08/96

Diln Fac: 1

LCS Lab ID: QC16520

Analyte	Result	Spike Added	%Rec #	Limits
Diesel Range	40.1	49.5	81	60-140
Surrogate	%Rec	Limits		
Hexacosane	69	60-140		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits

Curtis & Tompkins, Ltd.

Laboratory Number: 124634

Client: Parsons Engineering Science Inc.

Project#: 722450.14020 Location: Beale AFB Date Sampled: 02/28/96 Date Received: 02/29/96 Date Analyzed: 03/12/96

Batch: 26372

Analysis: Total Organic Carbon

Analysis Method: EPA 9060

LAB ID	SAMPLE ID	RESULT	UNITS	REPORTING	LIMIT
124634-004	BE-MP19-7.0	90	mg/Kg	50	
124634-Method	Blank	ND	mg/Kg	50	

ND = Not detected at or above reporting limit.

QA/QC SUMMARY:

MS/MSD RPD, %	<1	(Limit:	< 30)
MS/MSD AVG Recovery,%	87	(Limits:	50-150)
LCS Recovery,%	94	(Limits:	65-135)

QC'Sample: 124634-004



TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method:

LUFT

Sample # Cl	lient ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124634-001 BE	E-MP15-9.5	26268	02/27/96	03/05/96	03/09/96	10%
124634-002 BE	E-MP10-5.5	26268	02/28/96	03/05/96	03/08/96	10%
124634-003 BE	E-MP11-7.0	26268	02/28/96	03/05/96	03/08/96	12%
124634-004 BE	E-MP19-7.0	26268	02/28/96	03/05/96	03/08/96	9%

Analyte Diln Fac:	Units	124634-001 1	124634-002 1	124634-003 1	124634-004 1
JP5	mg/Kg	73 Y	110 Y	5.5YH	1.3z
Diesel Range	mg/Kg	110 Y	120 Y	10 Y	1.3z
Motor Oil Range	mg/Kg	<5.6	<5.6	<5.7	<5.5
Surrogate					
Hexacosane	%REC	92	92	100	94

Y: Sample exhibits fuel pattern which does not resemble standard

Z: Sample exhibits unknown single peak or peaks

H: Heavier hydrocarbons than indicated standard



Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 9471O, Phone (510) 486-0900

ANALYTICAL REPORT

Prepared for:

Parsons Engineering Science, Inc. 1301 Marina Village Parkway Suite 200 Alameda, CA 94501

Date: 18-MAR-96

Lab Job Number: 124634

Project ID: 722450.14020

Location: Beale AFB

Reviewed by:

Reviewed by:

This package may be reproduced only in its entirety.



Laboratory Number: 124634 Sample Date: 02/27,28/96 Client: Parsons Engineering Science Receipt Date: 02/29/96

Project#: 722450.14020 Location: Beale AFB

CASE NARRATIVE

Curtis & Tompkins Ltd. received four soil samples from the above referenced project on Feb.29,1996. All samples were received cold and intact. Chromatograms are included in this report for any samples containing quantifiable concentrations of extractable or purgeable hydrocarbons, along with example chromatograms for each type of fuel.

TOC (EPA 9060): No analytical problems were encountered.

TPH/Extractable: The "Z" qualifier flag appears for sample BE-MP19-7.0 (C&T# 124634-004) indicating that the chromatogram does not resemble a fuel pattern, but consists of individual, strongly differentiated peaks.

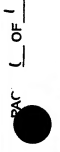
TPH/Purgeable: The chromatograms for these samples do not resemble gasoline, but are similar to a JP-4 standard which is occasionally requested. A JP-4 chromatogram is also included in this report for comparison purposes.

BTXE + TMBs (EPA 8020): Trifluorotoluene failed surrogate recovery limits for sample BE-MP10-5.5 (C&T# 124634-002) due to coelution of sample hydrocarbons with the surrogate. The "C" qualifier flag indicates that the target compound reported was confirmed by a second analytical column, but that the concentration quantitated on the second column differed from the reported concentration by more than a factor of two. This often happens when the fuel pattern present in the samples does not resemble gasoline, as is the case with this sample set. Ethylbenzene and xylene matrix spike and spike duplicate recoveries failed QC limits due to high levels of non-target hydrocarbons present in the spiked sample. The sample spiked was not from Beale AFB and the associated laboratory control sample passed all acceptance criteria.

PARSON'S ENGINEE SCIE

1301 Marina Village Pkwy, Suite 200 Alameda, California 94501 Phone: (510) 769-0100 FAX: (510) 769-9244





Project No.: 722450,14020	20				•				1	ORGANICS	SS			INORGANICS	NICS	ОТНЕЯ
Project Name/Location: Beale AFB, UST Site	Beale AFB,	UST Site	•					(2 1/2	, e'e		<u> </u>	\	-	1	////
Project Manager: Todd Wiedemeler (Denver, CO)	Wiedemeler	(Denver,	(00					201	\$610.	30	9701	\	<u> </u>	-	\	///
Site Manager: Michael Phelps (Alameda, CA)	helps (Alam	eda, CA)					•	NET!	00 (U	ov ee				1	(9)	
Sampler(s): (Initials and Signature(s)	itals and Signature(s)	12:				OF CONTA	SELITIVE	SET (JANF) SE (M2108/082E) H	0108/0E0S) \$70, \$ (0208/0E0S) \$5, \$ (0208/0E0S) \$5, \$ (08/13M70) } \$ (ws 108/0SSE) H. \$ \$ (ws 108/0E0S)	25 (5030/8020) 85 (0508/0503) 85 (05		(604 Abb) (605-3) 84 (6061)	(E.385.3)	ISS-O MISA) W	ISO MIST N	
Sample ID	Date	Time	Begin Depth	End	Matrix		SUNOSBB-H	of eldedos	MT brayo	(052E) \$0	Alink	(S.1357) Blinky (Stal & Nortosonal Le	2109) 401	etho Conta		
1 BE-M109-6	76H2h	0 \$40		\rightarrow	soil]-	A X	8 ×	<i>BY</i> − <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i> = <i>Y</i>	ο̈́ζ	200	वर्	40	20	T	Tu Az :
BE-MP12-4		1055	_	4.5	soil	_										ME O
3 BE-MP13-6,5	2/2496	0821		7.0	soil	_	X									= :
BE-MP 14-7	2/24/96 1315	1315	7.0	7.5	soil	-	X	×						17.7		" "
8 BE-4422-7	2/27/2	1510	7.0	7.5	soil	,	メメ	ス						783		11
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BELINQUISHED BY: (SIGNATURE)		E TIME		EIVED BY:	RECEIVED BY: (SIGNATUR	E)		NOTES:	1 20 20							
June 1		5						2. Quantif 3. Deloniz	2. Quantify all carbon ranges: diesel, jet fuel, oll/grease 3. Delonized water extraction	on ranges extraction	: dlesel,]	ot fuel, o	M/groese		- WD	4. I MiSK 1,2,3-timetry/ibenzene, 1,2,4- trimetry/ibenzene, and 1,3,5- trimetry/ibenzene.
NELINQUISHED BY: (SIGNATURE)	NTURE) DATE	TINE	-	EIVED FO	RECEIVED FOR LAB BY: (SIGNATURE)	SIGNAT		DATE	TIME	REMAR	KS/CC	DIMMENTS OF]]	L hork	3	
			D	2330	apullhung	12	45 7	7)			2		
Distribution: Orlainel acco	mpople of	1		:			1									

Pistribution: Original accompanies shipment; photocopy kept by samplers, copy FAXed to Parsons ES Site Manager by laboratory upon receipt of semples

Beale

COOLER RECEIPT CHECKLIST

Curtis	24	Tompkins,	110
	Ų,	TOTTOPRIES.	

	174133	(
Logina	#: 12162 Date Received: Number of Coolers:		
Client	: Parsu Fur. Junea Project: 722450.14020 Bea	ge	
	Parliminant Evanientian Phase		_
A.	Preliminary Examination Phase Date Opened: 2/28 By (print): (sign)	عريف	<u>Q</u> _
1.	Did cooler come with a shipping slip (airbill, etc.)? If YES, enter carrier name and airbill number: Feel 4 1154092418	(E)	NO
~	Were custody seals on outside of cooler?	YES	60
2.	We'le custody seats oil outside of cooler:		010
•	How many and where? Seal date: Seal name: Were custody seals unbroken and intact at the date and time of arrival?	VEC	KION
3.	Were custody seals unbroken and intact at the date and time of arrival	ILS	NO
4.	Were custody papers dry and intact when received?		
5.	Were custody papers filled out properly (ink, signed, etc.)?		NO.
6.	Did you sign the custody papers in the appropriate place?		
7.	Was project identifiable from custody papers?		МО
	If YES, enter project name at the top of this form.		110
8.	If required, was sufficient ice used? Type of ice: 3.75° L	E3	NO
	Type of ice: uhe ble Temperature: 3.75°C		
_	, -		
B.	Date Logged In: 2/21 By (print): \\ \(\text{U.II.s} \) (sign)	100	٦
	Date Logged In: 121 By (print): dw.llhs (sign)	<u>we</u>	
l.	Describe type of packing in cooler:		
2.	Did all bottles arrive unbroken?	" KES	NO
3.	Were labels in good condition and complete (ID, date, time, signature, etc.)?	12	
4.	Did bottle labels agree with custody papers?		NO
5.	Were appropriate containers used for the tests indicated?		NO
6.	Were correct preservatives added to samples?		NO
7.	Was sufficient amount of sample sent for tests indicated?	, ,	NO 1
8.	Were bubbles absent in VOA samples? If NO, list sample Ids below		
9.	Was the client contacted concerning this sample delivery?	YES	ио ,
	If YES, give details below.		
	Who was called? By whom? Dat	e:	
Additio	onal Comments:		÷
Filename:	F.\qc\forms\cooler.wpd Re	v. 1 4/95	

Filename: F.\qcvforms\cooler.wpd



BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Soil Batch#: 26389 Prep Date: Analysis Date: 03/13/96 03/13/96

Units: ug/Kg Diln Fac: 1

LCS Lab ID: QC16982

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	98	100	98	80-120
Toluene	100	100	100	80-120
Ethylbenzene	95	100	95	80-120
m,p-Xylenes	191	200	96	80-120
o-Xylene	102	100	102	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	87	43-114		
Bromobenzene	74	45-140		

Column to be used to flag recovery and RPD values with an asterisk alues outside of QC limits

ke Recovery: 0 out of 5 outside limits

BATCH QC REPORT

Page 1 of '

	BTX	E	
Project#:	Parsons Engineering Science, Inc. 722450.14020 Beale AFB	Analysis Method: Prep Method:	EPA 8020 EPA 5030
to grown gazer edite	MATRIX SPIKE/MATRIX	SPIKE DUPLICATE	× .*
Field ID: Lab ID: Matrix: Batch#: Units:	ZZZZZZ 124736-001 Soil 26345 ug/Kg dry weight	Sample Date: Received Date: Prep Date: Analysis Date: Moisture:	02/26/96 02/29/96 03/11/96 03/11/96 8%

MS Lab ID: QC16894

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Benzene	108.7	<5.435	109.8	101	75-125
Toluene	108.7	<5.435	105.4	97	75-125
Ethylbenzene	108.7	<5.435	83.7	77	75-125
m,p-Xylenes	217.4	<5.435	157.6	72 *	75-125
o-Xylene	108.7	<5.435	115.2	106	75-125
Surrogate	%Rec	Limits			· · · · · · · · · · · · · · · · · · ·
Trifluorotoluene	93	43-114			
Bromobenzene	69	45-140			

MSD Lab ID: QC16895

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Benzene	108.7	98.91	91	75-125	10	<20
Toluene	108.7	90.22	83	75-125	16	<20
Ethylbenzene	108.7	71.74	66 *	75-125	15	<20
m,p-Xylenes	217.4	132.6	61 *	75-125	17	<20
o-Xylene	108.7	108.7	100	75-125	6	<20
Surrogate	%Rec	Limit	8			
Trifluorotoluene	93	43-11	4			
Bromobenzene	70	45-14	0			

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits RPD: 0 out of 5 outside limits

Spike Recovery: 3 out of 10 outside limits



BATCH QC REPORT

Page 1 of 1

		·	BTXE		
Client: Project#: Location:	722450.		Inc.	Analysis Method: Prep Method:	
			METHOD BL	ANK	
Matrix: Batch#: Units: Diln Fac:	Soil 26389 ug/Kg 1			Prep Date: Analysis Date:	03/12/96 03/12/96

MB Lab ID: QC16980

Analyte	Result	
Benzene	<5.0	
Toluene	<5.0	
Ethylbenzene	<5.0	
m,p-Xylenes	<5.0	
o-Xylene	<5.0	
1,2,3-Trimethylbenzene	<5.0	
1,2,4-Trimethylbenzene	<5.0	
1,3,5-Trimethylbenzene	<5.0	
Surrogate	%Rec	Recovery Limits

Surrogate	%Rec	Recovery Limits
rifluorotoluene	94	43-114
komobenzene	82	45-140



BATCH QC REPORT

Page 1 of !

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Soil

Batch#: 26345 Units: ug/Kg Diln Fac: 1 Prep Date: 03/11/96 Analysis Date: 03/11/96

LCS Lab ID: QC16893

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	106	100	106	80-120
Toluene	109	100	109	80-120
Ethylbenzene	105	100	105	80-120
m,p-Xylenes	214	200	107	80-120
o-Xylene	109	100	109	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	89	43-114		
Bromobenzene	74	45-140		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 5 outside limits



Page 2 of 2

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample # Client I	Batch #	Sampled	Extracted	Analyzed	Moisture
124633-005 BE-MP22-	26389	02/27/96	03/12/96	03/12/96	20%

Analyte	Units	124633-	005	•
Diln Fac:		25		•
Benzene	ug/Kg	<16		
Toluene	ug/Kg	<16		
Ethylbenzene	ug/Kg	<16		
m,p-Xylenes	ug/Kg	150	С	
o-Xylene	ug/Kg	<16		
1,2,3-Trimethylbenz	ene ug/Kg	78		
1,2,4-Trimethylbenz	ene ug/Kg	70	С	
1,3,5-Trimethylbenz	ene ug/Kg	36		
Surrogate				
Trifluorotoluene	%REC	94		
Bromobenzene	%REC	78		

C: Presence of this compound confirmed by second column, however, the confirmation concentration differed from the reported result by more than a factor of two



BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

METHOD BLANK

Matrix: Soil Batch#:

Diln Fac: 1

26345

Units: ug/Kg

Prep Date: 03/11/96 Analysis Date:

03/11/96

MB Lab ID: QC16891

Analyte	Result	
Benzene	<5.0	
Toluene	<5.0	
Ethylbenzene	<5.0	
m,p-Xylenes	<5.0	
o-Xylene	<5.0	
1,2,3-Trimethylbenzene	<5.0	
1,2,4-Trimethylbenzene	<5.0	
1,3,5-Trimethylbenzene	<5.0	
Surrogate	%Rec	Recovery Limits

Surrogate	%Rec	Recovery Limits
Trifluorotoluene	87	43-114
Bromobenzene	71	45-140

: G:\GC04\B\026J011.raw Date: 2/1/95 3:57 PM Page 1 of 1 End Time : 17.00 min Start Time : 0.00 min Low Point : -4.15 mV High Point : 170.85 mV Scale Factor: -1 Plot Offset: -4 mV Plot Scale: 175 mV Response [mV] 0.93 -1.25 -1.46 -1.74 2.01 2.14 2,.33 TRIFLUOROT -2.66 3.06 TOLUENE -3.62 -3.81 4.02 -4.38 -5.06 .28 M, HYLYBENNE = 5.44 5.76.57 5.9785 BROMOBENZE -GAS PEAK 1 = GAS PEAK 3 -7.65 -7.93 10.02

Ret_.ition

Time



BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124633-001	BE-MP09-6	26345	02/27/96	03/12/96	03/12/96	14%
124633-002	BE-MP12-4	26345	02/27/96	03/12/96	03/12/96	16%
124633-003	BE-MP13-6.5	26345	02/27/96	03/12/96	03/12/96	14%
124633-004	BE-MP14-7	26345	02/27/96	03/12/96	03/12/96	15%

Analyte	Units	124633-001	124633-002	124633-003	124633-004
Diln Fac:		1	1	1	1
Benzene	ug/Kg	<5.8	<6	<5.8	<5.9
Toluene	ug/Kg	<5.8	<6	<5.8	<5.9
Ethylbenzene	ug/Kg	<5.8	<6	<5.8	<5.9
m,p-Xylenes	ug/Kg	<5.8	<6	<5.8	81 C
o-Xylene	ug/Kg	<5.8	<6	<5.8	<5.9
1,2,3-Trimethylben:	zene ug/Kg	<5.8	<6	<5.8	120
1,2,4-Trimethylben:	zene ug/Kg	<5.8	<6	<5.8	160
1,3,5-Trimethylben:	zene ug/Kg	<5.8	<6	<5.8	130
Surrogate					
Trifluorotoluene	%REC	86	88	64	133 *
Bromobenzene	%REC	75	76	53	76

^{*} Values outside of QC limits

C: Presence of this compound confirmed by second column, however, the confirmation concentration differed from the reported result by more than a factor of two

FileName : G:\GC04\072J006.raw Start Time : 0.00 min Er

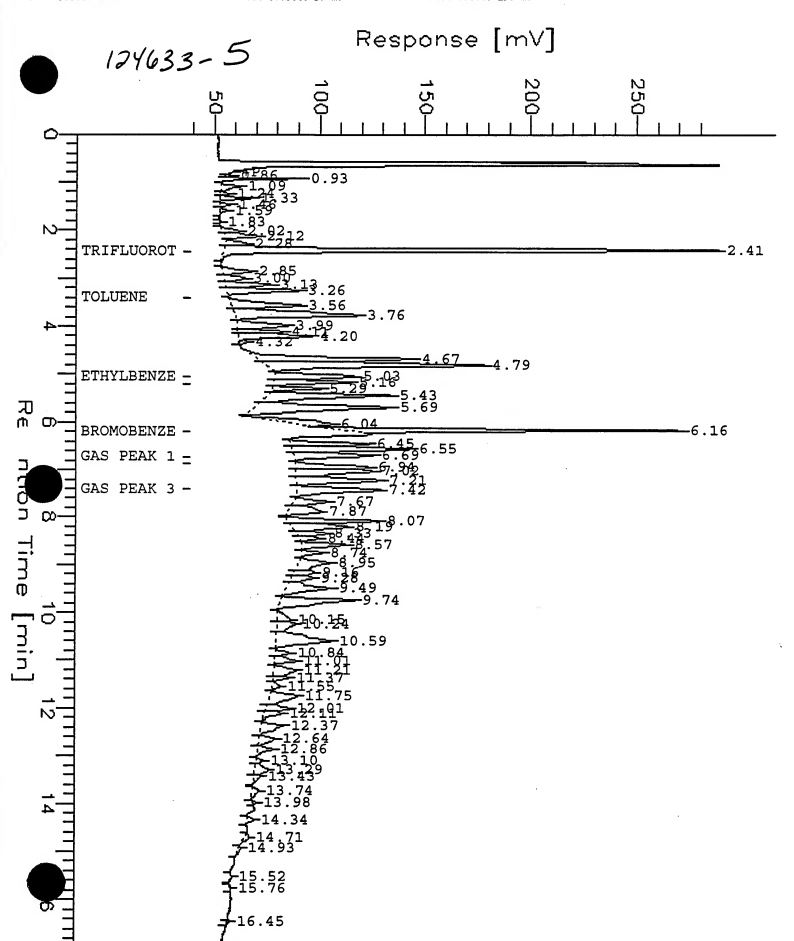
Scale Factor: -1

End Time : 17.00 min Plot Offset: 39 mV

Date : 3/12/96 4:56 PM Low Point : 38.90 mV

Plot Scale: 250 mV

Page 1 of 1 High Point: 288.90 mV



TVH2 - GC-04 RTX-1

C+T Gasoline Std. : G:\GC04\135J001.raw FileName

Start Time : 0.00 min

Time

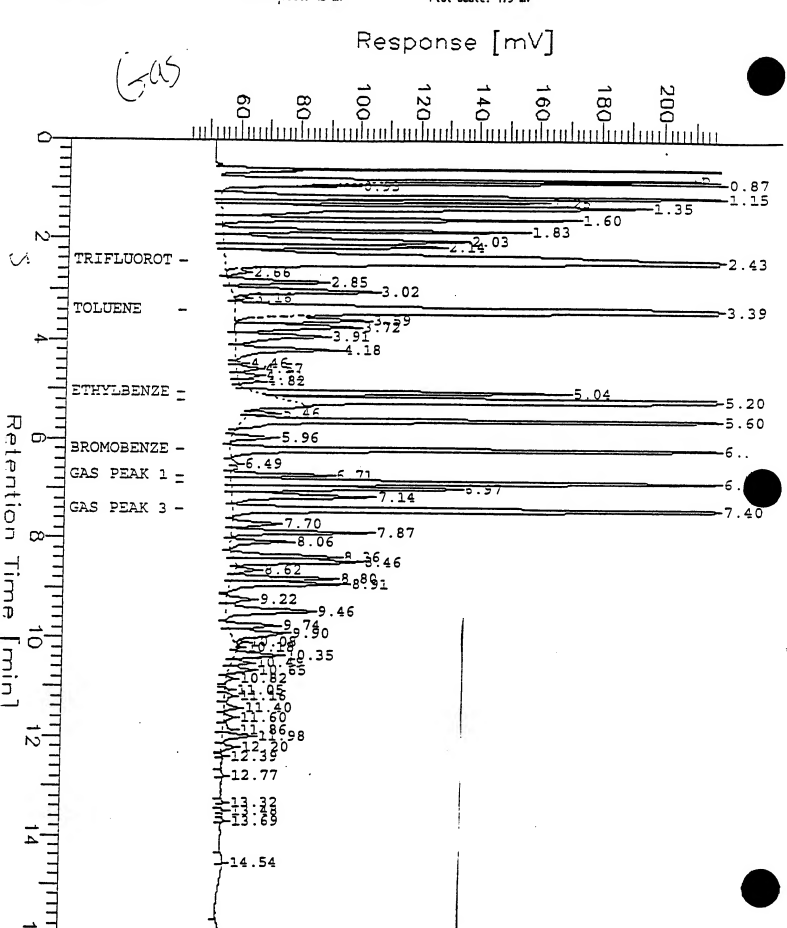
Scale Factor: -1

End Time : 17.00 min Plot Offset: 43 mV

16.10 -16.45 Date: 5/15/95 9:53 AM Low Point: 42.92 mV

Plot Scale: 175 mV

Page 1 of 1 High Point : 217.92 mV



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

F

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method:

EPA 5030

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Sample Date:

02/29/96

Field ID: ZZZZZZ Lab ID: 124660-002

Received Date:

03/04/96

Matrix: Soil Batch#: 26389

Prep Date:

03/13/96

Units: mg/Kg Diln Fac: 1 Analysis Date:

03/13/96

MS Lab ID: QC16983

Analyte	Spike Added	Sample	Sample MS		Limits	
Gasoline	10	<1.000	7.7	77	75-125	
Surrogate	%Rec	Limits				
Trifluorotoluene Bromobenzene	95 90	52-127 47-112				

D Lab ID: QC16984

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Gasoline	10	7.8	78	75-125	1	<20
Surrogate	%Rec	Limit	s			
Trifluorotoluene Bromobenzene	99 91	52-12 47-11	-			

[#] Column to be used to flag recovery and RPD values with an asterisk

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits RPD: 0 out of 1 outside limits

: G:\GC04\071J027.raw

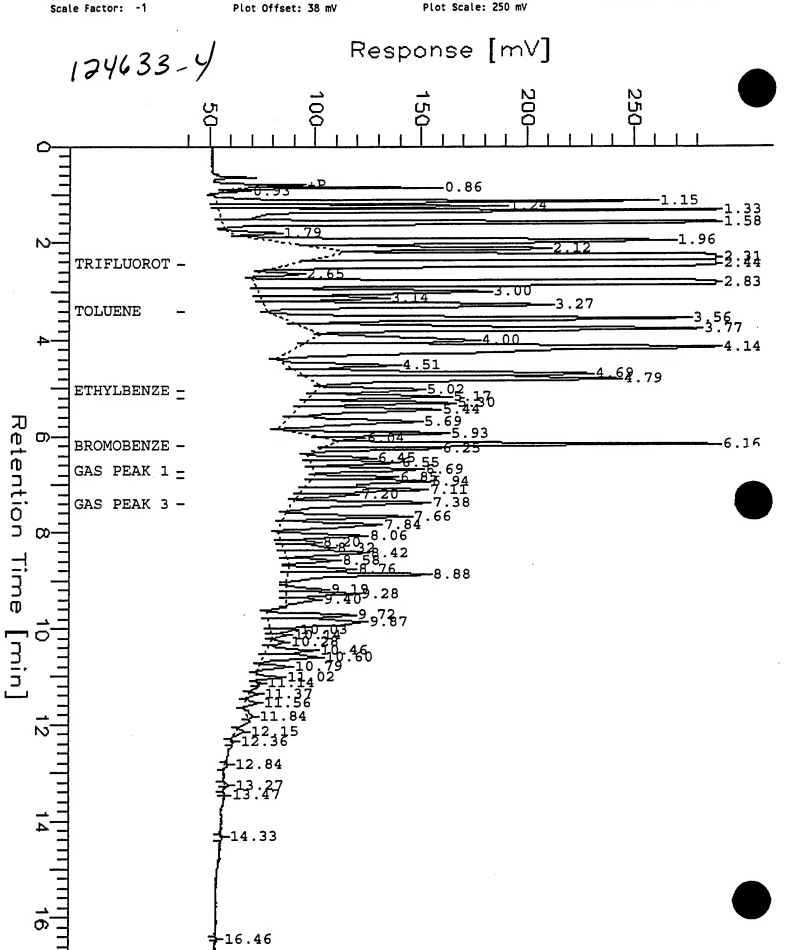
Start Time : 0.00 min

End Time : 17.00 min Plot Offset: 38 mV

Date: 3/12/96 2:25 AM Low Point: 38.41 mV

Plot Scale: 250 mV

Page 1 of 1 High Point: 288.41 mV



BATCH QC REPORT

Page 1 of 1

" in the original

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Location: Beale AFB

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Soil Prep Date:

03/11/96

Batch#: 26345 Units: mg/Kg Diln Fac: 1

Analysis Date: 03/11/96

LCS Lab ID: QC16892

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	10.1	10	101	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	106 90	52-127 47-112		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits 'pike Recovery: 0 out of 1 outside limits



03/12/96

03/12/96

Lab #: 124633

BATCH QC REPORT

Page 1 of '

TVH-Total Volatile Hydrocarbons

Parsons Engineering Science, Inc. Client:

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Soil Batch#: 26389 Prep Date: Analysis Date:

Units: mg/Kg Diln Fac: 1

LCS Lab ID: QC16981

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	10.2	10	102	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	108 89	52-127 47-112		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits

BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

roject#: 722450.14020 Prep Method:

EPA 5030

Location: Beale AFB

Soil

26345

METHOD BLANK

Prep Date: 03/11/96

Units: mg/Kg Analysis Date:

03/11/96

Diln Fac: 1

Matrix:

Batch#:

MB Lab ID: QC16891

Analyte	Result	
Gasoline	<1.0	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene Bromobenzene	93 78	52-127 47-112



BATCH QC REPORT

Page 1 of

TVH-Total Volatile Hydrocarbons

Parsons Engineering Science, Inc. Client:

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Location: Beale AFB

EPA 5030

Prep Method:

METHOD BLANK

Matrix: Soil

Prep Date:

03/12/96

Batch#: 26389 Units: mg/Kg Analysis Date:

03/12/96

Diln Fac: 1

MB Lab ID: QC16980

Analyte	Result	
Gasoline	<1.0	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene Bromobenzene	96 85	52-127 47-112



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Prep Date: 03/12/96 Analysis Date: 03/12/96

Matrix: Soil
Batch#: 26389
Units: mg/Kg
Diln Fac: 1

LCS Lab ID: QC16981

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	10.2	10	102	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	108 89	52-127 47-112		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits



BATCH QC REPORT

Page 1 of J

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Method: EPA 5030

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Sample Date: 02/29/96
Lab ID: 124660-002 Received Date: 03/04/96
Matrix: Soil Prep Date: 03/13/96
Batch#: 26389 Analysis Date: 03/13/96

Units: mg/Kg

Diln Fac: 1

MS Lab ID: QC16983

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Gasoline	10	<1.000	7.7	77	75-125
Surrogate	%Rec	Limits			· · · · · · · · · · · · · · · · · · ·
Trifluorotoluene Bromobenzene	95 90	52-127 47-112			

MSD Lab ID: QC16984

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Gasoline	10	7.8	78	75-125	1	<20
Surrogate	%Rec	Limit	:8			
Trifluorotoluene Bromobenzene	99 91	52-12 47-11	-			· · · · · · · · · · · · · · · · · · ·

[#] Column to be used to flag recovery and RPD values with an asterisk

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits RPD: 0 out of 1 outside limits

FileName : G:\GC04\072J005.raw
Start Time : 0.00 min E
Scale Factor: -1 P

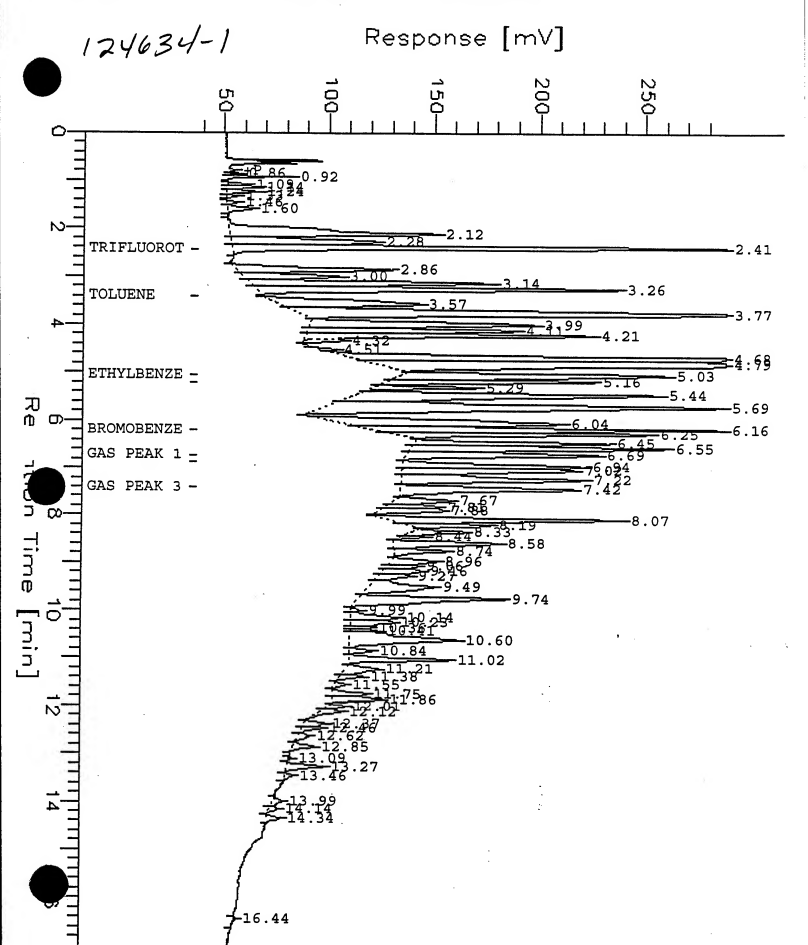
End Time : 17.00 min

Plot Offset: 38 mV

Date : 3/12/96 4:18 PM Low Point : 38.06 mV

Plot Scale: 250 mV

Page 1 of 1 High Point: 288.06 mV



FileName : G:\GC04\072J008.raw

Start Time : 0.00 min

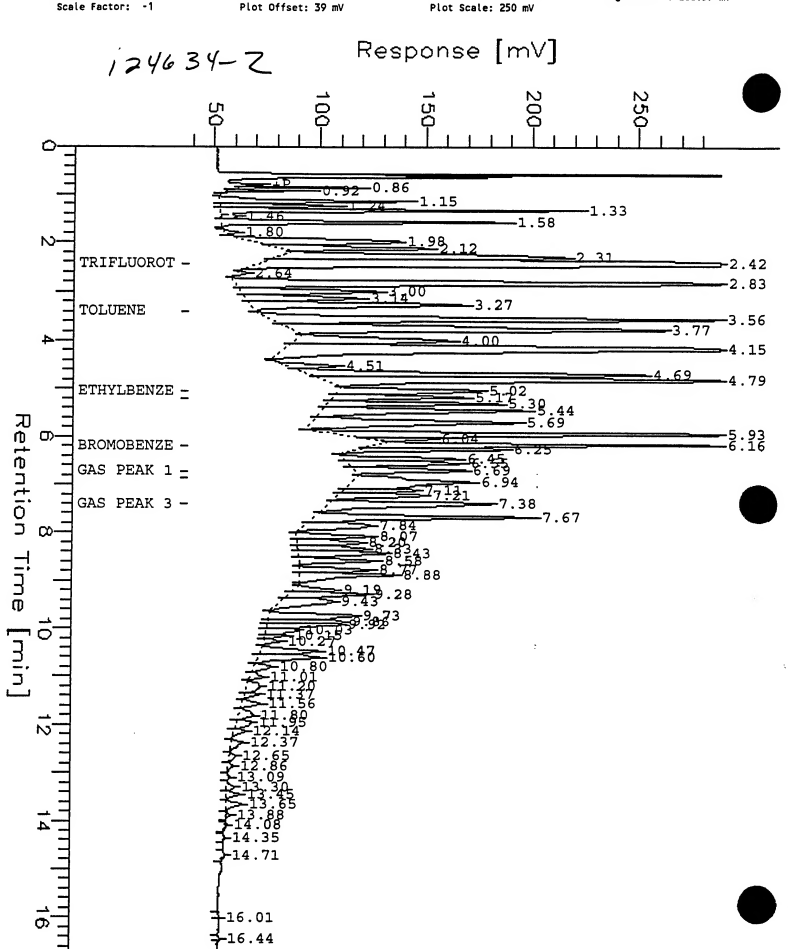
End Time : 17.00 min

Plot Offset: 39 mV

Date : 3/12/96 5:52 PM Low Point : 38.59 mV

Plot Scale: 250 mV

Page 1 of 1 High Point: 288.59 mV



FileName : G:\GC04\072J007.raw
Start Time : 0.00 min Er

End Time : 17.00 min

Date : 3/12/96 5:24 PM Low Point : 38.82 mV

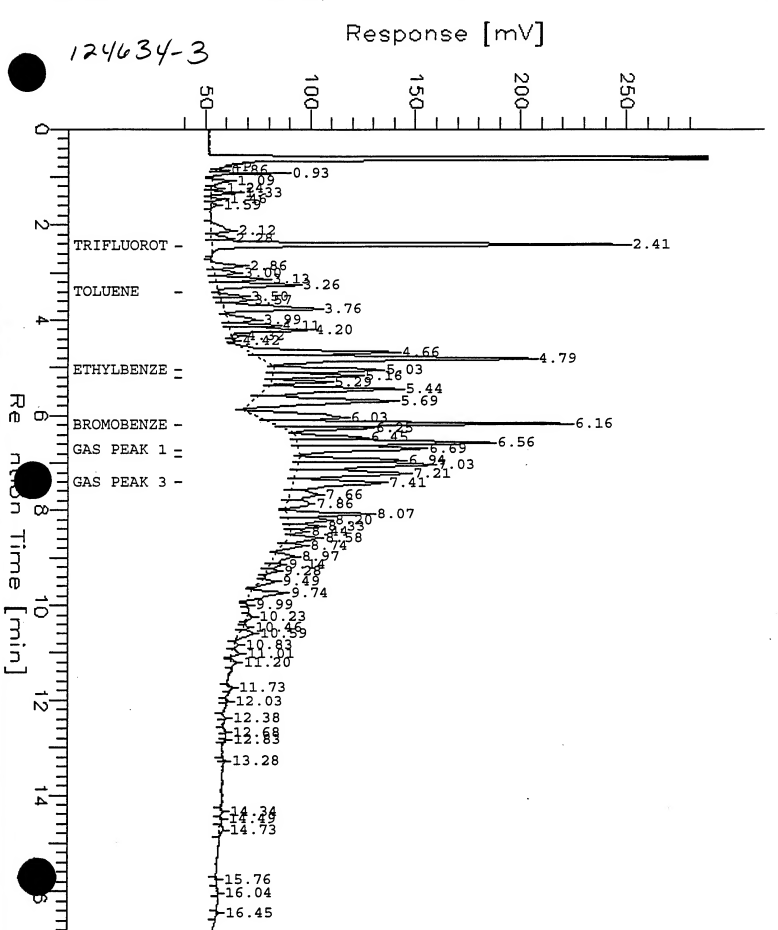
Plot Scale: 250 mV

Page 1 of 1

Scale Factor: -1

Plot Offset: 39 mV

High Point: 288.82 mV



C+T Gasoline Std. : G:\GC04\135J001.raw FileName Date : 5/15/95 9:53 AM Low Point : 42.92 mV Page 1 of 1 Start Time : 0.00 min End Time : 17.00 min High Point : 217.92 mV Scale Factor: -1 Plot Offset: 43 mV Plot Scale: 175 mV Response [mV] -0.87 1.15 1.35 2.03 TRIFLUOROT --2.43 2.85 -3.02 TOLUENE -3.39 3.91· 4.18 ETHYLBENZE = 5.04 -5.20 Relention 5.50 5.96 BROMOBENZE -6.] 5.49 GAS PEAK 1 = 7.14 GAS PEAK 3 -7.70 -7.87 8.06 - કે6 4 6 Time 62 9 74 0 9 74 0 10.40 0.35 11.40 11.40 11.40 -9.46 128698 12.390 12.77 13.32 13.69 -14.54 16.10 -16.45

: G:\GC04\B\026J011.raw Date: 2/1/95 3:57 PM Page 1 of 1 End Time : 17.00 min High Point: 170.85 mV Start Time : 0.00 min Low Point : -4.15 mV Plot Scale: 175 mV Scale Factor: -1 Plot Offset: -4 mV Response [mV] 0.93 تون خز. -1.25 1.46 -1.74 01 -2.14 TRIFLUOROT -2.66 -3.06 TOLUENE -3.62 $\frac{-3.81}{4.02}$ 4.38 4.58 -5.44 70 . 5 / 97 85 6.20 BROMOBENZE -6.44 GAS PEAK 1 = GAS PEAK 3 -7.65 -7.93 Time 10.02



BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124634-001 BE-MP15-9.5	26389	02/27/96	03/12/96	03/12/96	10%
124634-002 BE-MP10-5.5	26389	02/28/96	03/12/96	03/12/96	10%
124634-003 BE-MP11-7.0	26389	02/28/96	03/12/96	03/12/96	12%
124634-004 BE-MP19-7.0	26345	02/28/96	03/12/96	03/12/96	9%

Analyte Diln Fac:	Units	124634-001 1	124634- 200	002	124634- 25	-003	124634-004 1
Benzene	ug/Kg	<5.6	<110		<14		<5.5
Toluene	ug/Kg	<5.6	<110		<14		<5.5
Ethylbenzene	ug/Kg	<5.6	<110		<14		<5.5
m,p-Xylenes	ug/Kg	86 C	2200	С	160	С	<5.5
o-Xylene	ug/Kg	<5.6	<110		<14		<5.5
1,2,3-Trimethylbenz	ene ug/Kg	190	<110		110		<5.5
1,2,4-Trimethylbenz	ene ug/Kg	<5.6	180	С	85	С	<5.5
1,3,5-Trimethylbenz	ene ug/Kg	90	<110		16	С	<5.5
Surrogate							
Trifluorotoluene	%REC	100	135	*	67		84
Bromobenzene	%REC	77	77		56		72

^{*} Values outside of QC limits

C: Presence of this compound confirmed by second column, however, the confirmation concentration differed from the reported result by more than a factor of two



BATCH QC REPORT

Page 1 of 1

BTXE

Parsons Engineering Science, Inc.

Client: Parsons Engineroject#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

METHOD BLANK

Matrix: Soil Batch#: 26345

Units: ug/Kg Diln Fac: 1

Prep Date: 03/11/96 Analysis Date: 03/11/96

MB Lab ID: QC16891

Analyte	Result	
Benzene	<5.0	
Toluene	<5.0	
Ethylbenzene	<5.0	
m,p-Xylenes	<5.0	
o-Xylene	<5.0	
1,2,3-Trimethylbenzene	<5.0	
1,2,4-Trimethylbenzene	<5.0	
1,3,5-Trimethylbenzene	<5.0	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene .	87	43-114
romobenzene	71	45-140



BATCH QC REPORT

Page 1 of 1

BTXE							
Project#:	Parsons Engineering Scienc 722450.14020 Beale AFB	e, Inc.	Analysis Method: Prep Method:	EPA 8020 EPA 5030			
		METHOD	BLANK				
Matrix: Batch#: Units: Diln Fac:	Soil 26389 ug/Kg 1		Prep Date: Analysis Date:	03/12/96 03/12/96			

MB Lab ID: QC16980

Analyte	Result		
Benzene	<5.0		
Toluene	<5.0		
Ethylbenzene	<5.0		
m,p-Xylenes	<5.0		
o-Xylene	<5.0		
1,2,3-Trimethylbenzene	<5.0		
1,2,4-Trimethylbenzene	<5.0		
1,3,5-Trimethylbenzene	<5.0		
Surrogate	%Rec	Recovery Limits	
Trifluorotoluene	94		
Bromobenzene	82	45-140	

BATCH QC REPORT

Page 1 of 1

BTXE

lient: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

Project#: 722450.14020 Location: Beale AFB Prep Method: EPA 5030

Prep Method

LABORATORY CONTROL SAMPLE

 Matrix:
 Soil
 Prep Date:
 03/11/96

 Batch#:
 26345
 Analysis Date:
 03/11/96

Units: ug/Kg
Diln Fac: 1

LCS Lab ID: QC16893

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	106	100	106	80-120
Toluene	109	100	109	80-120
Ethylbenzene	105	100	105	80-120
m,p-Xylenes	214	200	107	80-120
o-Xylene	109	100	109	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	. 89	43-114		
Bromobenzene	74	45-140		
Bromoberizene	/4	45-140		

Column to be used to flag recovery and RPD values with an asterisk alues outside of QC limits

ke Recovery: 0 out of 5 outside limits



Page 1 of 1

Lab #: 124634 BATCH QC REPORT

BTXE

Parsons Engineering Science, Inc. Client:

Project#: 722450.14020

Soil

26389

Analysis Method: EPA 8020

Prep Method:

EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Prep Date:

03/13/96

Analysis Date: 03/13/96

Units: ug/Kg Diln Fac: 1

Matrix:

Batch#:

LCS Lab ID: QC16982

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	98	100	98	80-120
Toluene	100	100	100	80-120
Ethylbenzene	95	100	95	80-120
m,p-Xylenes	191	200	96	80-120
o-Xylene	102	100	102	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	87	43-114		
Bromobenzene	74	45-140		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 5 outside limits

BATCH QC REPORT

Page 1 of 1

BTXE

lient: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Lab ID: 124736-001

Matrix: Soil Batch#: 26345

Units: ug/Kg dry weight

Diln Fac: 1

Sample Date: 02/26/96 Received Date: 02/29/96

Prep Date: 03/11/96 Analysis Date: 03/11/96

Moisture: 88

MS Lab ID: QC16894

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Benzene	108.7	<5.435	109.8	101	75-125
Toluene	108.7	<5.435	105.4	97	75-125
Ethylbenzene	108.7	<5.435	83.7	77	75-125
m,p-Xylenes	217.4	<5.435	157.6	72 *	75-125
o-Xylene	108.7	<5.435	115.2	106	75-125
Surrogate	%Rec	Limits			
Trifluorotoluene	93	43-114			· · · · · · · · · · · · · · · · · · ·
Bromobenzene	69	45-140			

MSD Lab ID: QC16895

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Benzene	108.7	98.91	91	75-125	10	<20
Toluene	108.7	90.22	83	75-125	16	<20
Ethylbenzene	108.7	71.74	66 *	75-125	15	<20
m,p-Xylenes	217.4	132.6	61 *	75-125	17	<20
o-Xylene	108.7	108.7	100	75-125	6	<20
Surrogate	%Rec	Limit	s			
Trifluorotoluene	93	43-11	4			
Bromobenzene	70	45-14	-			

[#] Column to be used to flag recovery and RPD values with an asterisk

RPD: 0 out of 5 outside limits

Spike Recovery: 3 out of 10 outside limits

^{*} Values outside of QC limits

PARSONS ENGINEERING SCIENCE, INC.

1301 Marina Village Pkwy, Suite 200 Alameda, California 94501

Phone: (510) 769-0100 FAX: (510) 769-9244

CHAIN OF CUSTODY RECORD 124001

OF |

PAGE \

handbook requirements. 4. TMBs: 1,2,3-trimethylbenzene, 1,2,4-trimethylbenzene, and 1,3,5-trimethylbenzene. according to AFCEE and holding times Note: All enalyses Roth Remarks , î OTHER THE NEDWA TMBs RACHWAD BY. ٦ ٤ Site Manager INORGANICS Secondary column confirmation required.
 Quantify all carbon ranges: diesel, jet fuel, oll/grease.
 Delonized water extraction REMARKS/COMMENTS: cheek in. 6100 663 (0108/0E02) ORGANICS 18 15 ES TIME X ANALYTES (METHOD) NOTES: RECEIVED FOR LAB BY: (SIGNATURE) DATE X 义 人 Marcus J. Prein NO. OF CONTAINERS RECEIVED BY: (SIGNATURE) Matrix Sol Soil Soll soi Soll <u>8</u> SO. SO SO <u>8</u>0 <u>8</u> 80 8 Depth 00 0.0 7.5 7.5 Begin Depth 150 r Di 95 4.0 Project Manager: Todd Wiedemeier (Denver, CO) 2/22/20 1543 2/4/96/11 35 BE-HP19-7.0 2/28/96 1640 Project Name/Location: Beale AFB, UST Site 150 Site Manager: Michael Phelps (Alameda, CA) 1130 Time 78 DATE DATE BE-MP11-70 428/96 Sampler(s): (Initials and Signature(s)) BE-MP19-55/4/28/96 Marine J. 96/22/2 Date BY: (SIGNATURE) RELINQUISHED BY (SIGNATURE) Project No.: 722450.14020 BE-MPS-45 MED Sample ID RELINGUISHED

tribution: Original accompanies shipment; photocopy kept by samplers; copy FAXed to Parsons ES Site Manager by laboratory upon receipt of samples.

COOLER RECEIPT CHECKLIST



	#: 124634 Date Received: 2(21)	Number of Cooler	(
Logini	Date Received:	Number of Cooler 722450.	
Client	rioject. Ma	Ju 10000	19020
A. ·	Preliminary Examination Phase Date Opened: 22 By (print): 50	(Sign)	Gusée-
1.	Did cooler come with a shipping slip (airbill, etc. If YES, enter carrier name and airbill number:	c.)?	YES (NO
2.	Were custody seals on outside of cooler?		YES NO
3.	How many and where? Seal Were custody seals unbroken and intact at the	iate and time of arrival?.	YES NO
4.	Were custody papers dry and intact when receive	ved?	
5.	Were custody papers filled out properly (ink, si	gned, etc.)?	ON SEC
6.	Did you sign the custody papers in the appropri	ate place?	
7.	Was project identifiable from custody papers?	•	MED NO
	If YES, enter project name at the top of this for		
3.	If required was sufficient ice used?		YES NO
	Type of ice: Luke blue Tem	perature: 4,0°C	
	7	_	
B.	Login Phase Date Logged In: 2/24 By (print): 2.W.	(sign) →	- هدن هم
1.	Describe type of packing in cooler:		
2.	Did all bottles arrive unbroken?		XES NO
3.	Were labels in good condition and complete (II), date, time, signature, e	tc.)?
4.	Did bottle labels agree with custody papers?		
5.	Were appropriate containers used for the tests i	ndicated?	NO
6.	Were correct preservatives added to samples?		NO NO
7.	Was sufficient amount of sample sent for tests i	ndicated?	NO SES NO
3.	Were bubbles absent in VOA samples? If NO, li	ist sample Ids below	YES NO んり
9.	Was the client contacted concerning this sample		
	If YES, give details below.	•	
	Who was called? By w	vhom?	Date:
Additio	ional Comments:		:
Filename	e: F.\qc\forms\cooler.wpd		Rev. 1 4/95



Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 9471O, Phone (510) 486-0900

ANALYTICAL REPORT

Prepared for:

Parsons Engineering Science, Inc. 1301 Marina Village Parkway Suite 200 Alameda, CA 94501

Date: 21-MAR-96

Lab Job Number: 124647

Project ID: 722450.14020

Location: Beale AFB

Reviewed by: Just K Morriso

Reviewed by:

This package may be reproduced only in its entirety.



Laboratory Number: 124647

Client: Parsons Engineering Science

Project#: 722450.14020 Location: Beale AFB Sample Date: 03/01/96 Receipt Date: 03/02/96

CASE NARRATIVE

Curtis & Tompkins Ltd. received three soil samples from the above referenced project on March 1,1996. All samples were received cold and intact. The chain of custody requested matrix spike and spike duplicate on sample BE-MP01-5 (C&T# 124647-002), however, a large fraction of each of these samples were large pebbles and rocks. There was not enough soil to perform spikes on any of the samples in this set. Chromatograms are included in this report for any samples containing quantifiable concentrations of extractable or purgeable hydrocarbons, along with example chromatograms for each type of fuel.

TPH/Extractable: The "Z" qualifier flag appears for sample BE-MP23-6.5 (C&T# 124647-001) indicating that the sample chromatogram does not resemble a fuel pattern, but consists of individual, strongly differentiated peaks.

TPH/Purgeable: No analytical problems were encountered.

BTXE + TMBs (EPA 8020): Ethyl-benzene and xylene matrix spike and spike duplicate recoveries failed QC limits due to high levels of non-target hydrocarbons present in the spiked sample. The sample spiked was not from Beale AFB and the associated laboratory control sample passed all acceptance criteria.



TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc.

roject#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: LUFT

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124647-001	BE-MP23-6.5	26268	03/01/96	03/05/96	03/08/96	12%
124647-002	BE-MP01-5	26268	03/01/96	03/05/96	03/09/96	12%
124647-003	BE-MP43-7.0	26268	03/01/96	03/05/96	03/08/96	13%

Analyte Diln Fac:	Units	124647-001 1	124647-002 1	124647-003 1	
JP5	mg/Kg	1.9Z	<1.1	<1.1	
Diesel Range	mg/Kg	<1.1	4.4YH	<1.1	
Motor Oil Range	mg/Kg	<5.7	51	<5.7	
Surrogate					
"exacosane	%REC	107	92	97	

Sample exhibits fuel pattern which does not resemble standard

Z: Sample exhibits unknown single peak or peaks

H: Heavier hydrocarbons than indicated standard



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

thaman 1 Mar I is

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method:

Location: Beale AFB

METHOD BLANK

Matrix: Soil Batch#: 26268 Prep Date: Analysis Date: 03/08/96

03/05/96

Units: mg/Kg Diln Fac: 1

MB	Lab	ID:	ÕC19	519

Analyte	Result	
JP5	<1.0	
Diesel Range	<1.0	
Motor Oil Range	<5.0	
Surrogate	%Rec	Recovery Limits
Hexacosane	69	60-140



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: LUFT

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Prep Date:

03/05/96

Matrix: Soil Batch#: 26268

Analysis Date: 03/08/96

Units: mg/Kg Diln Fac: 1

LCS Lab ID: QC16520

Analyte	Result	Spike Added	%Rec #	Limits
Diesel Range	40.1	49.5	81	60-140
Surrogate	%Rec	Limits		
Hexacosane	69	60-140		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits

BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: LUFT

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Lab ID: 124641-004

Matrix: Soil Batch#: 26268

Units: mg/Kg dry weight

Diln Fac: 1

Sample Date:

02/27/96 Received Date: 03/01/96 Prep Date:

03/05/96 Analysis Date: 03/08/96

Moisture: 35%

MS Lab ID: QC16521

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Diesel Range	76.15	<1.538	71.69	94	60-140
Surrogate	%Rec	Limits			
Hexacosane	82	60-140			

MSD Lab ID: QC16522

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Diesel Range	76.15	63.54	83	60-140	12	<30
Surrogate	%Rec	Limits				7
Hexacosane	74	60-14	0			

[#] Column to be used to flag recovery and RPD values with an asterisk

RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits

GC15 Channel A Surrogate

Sample Name : S,124647-001,26268

FileName : C:\GC15\CHB\068B009.raw

Method : DUAL Start Time : 0.00 min

End Time : 31.90 min

Factor: 0.0 Plot Offset: 32 mV

Sample #: 26268

Page 1 of 1

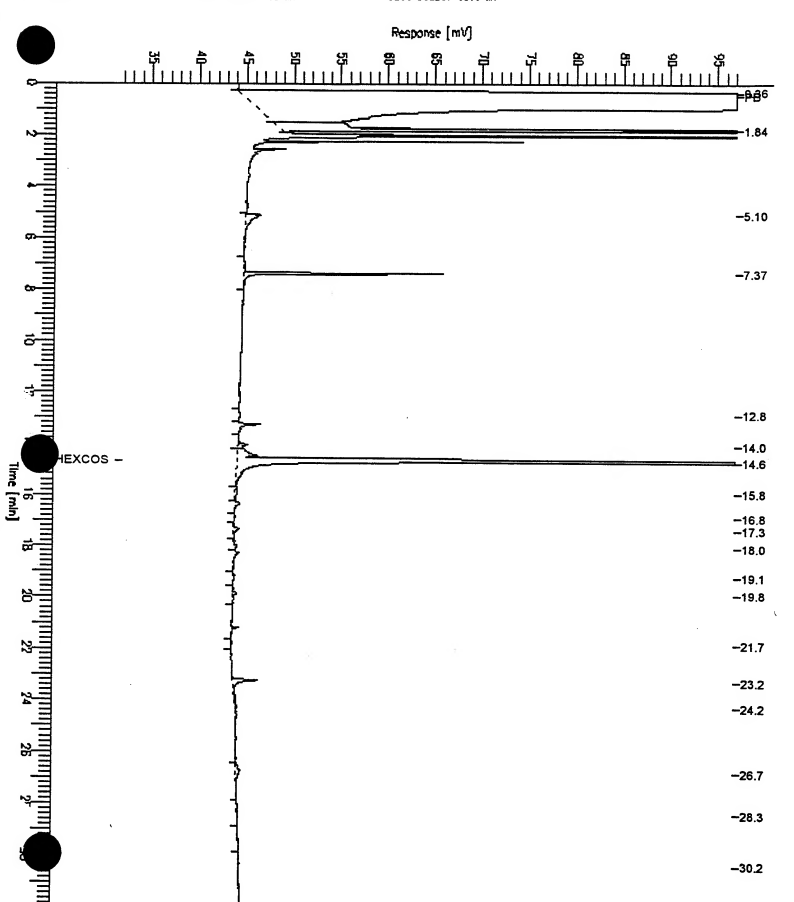
Date : 3/8/96 11:26 PM

Time of Injection: 3/8/96 10:53 PM

Low Point : 32.00 mV

High Point : 97.00 mV

Plot Scale: 65.0 mV



Sample Name: S,124647-002,26268

FileName : C:\GC15\CHB\068B022.raw

Method : DUAL

Start Time : 0.00 min

End Time : 31.90 min

Plot Offset: 32 mV

Sample #: 26268

Date: 3/9/96 08:55 AM

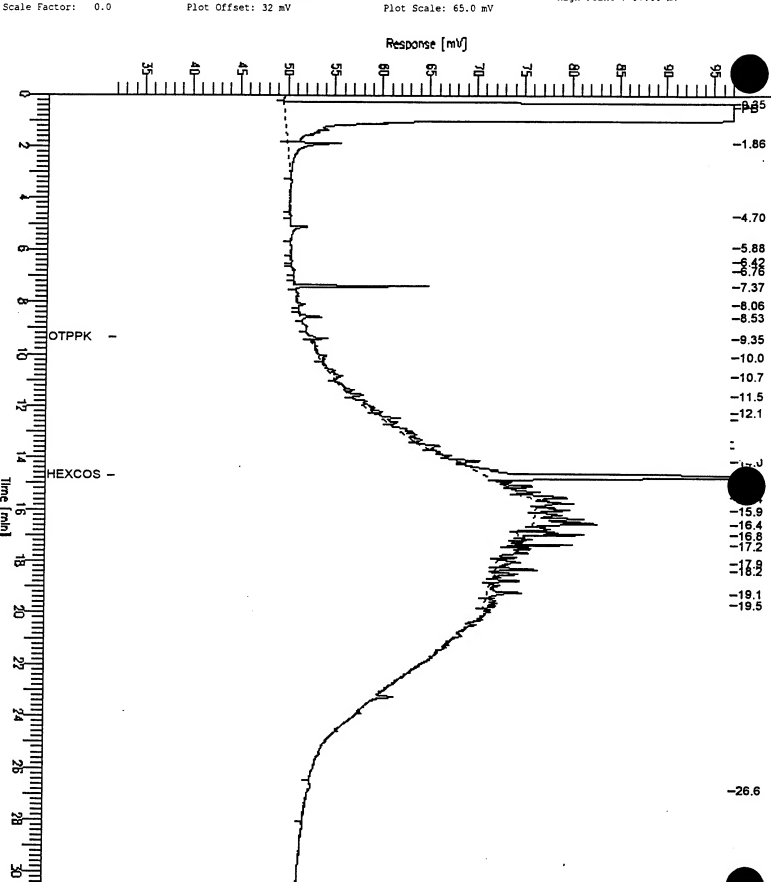
Time of Injection: 3/9/96 08:22 AM

Low Point : 32.00 mV

High Point: 97.00 mV

Page 1 of 1

Plot Scale: 65.0 mV



JP-5

GC15 Channel A TEH

Sample Name : CCV, JP5

: C:\GC15\CHB\067B023.RAW FileName

: BTEH.MTH Method Start Time : 0.01 min

End Time

Factor: 0.0

: 26.87 min

Plot Offset: 42 mV

Sample #: 250MG/L

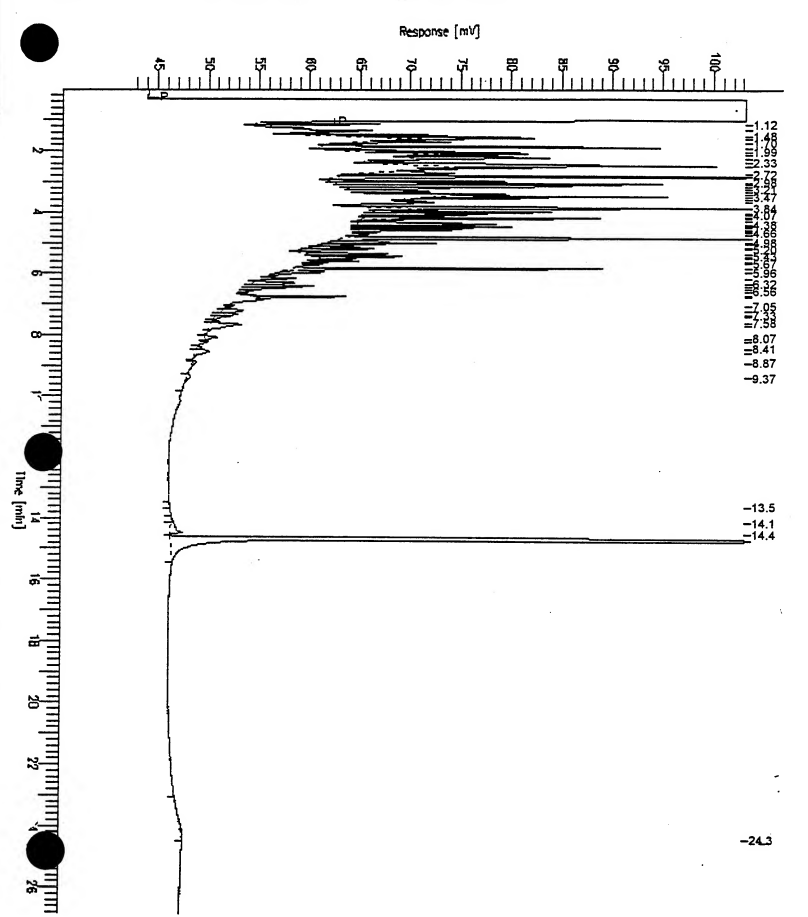
Date: 3/8/96 01:50 PM

Time of Injection: 3/8/96 07:20 AM Low Point: 42.42 mV

Plot Scale: 60.8 mV

High Point : 103.22 mV

Page 1 of 1



Clidillier A GCID TU

Sample Name : CCV, 96WS1948, DSL

: C:\GC15\CHB\067B021.RAW FileName

: BTEH.MTH Method

Start Time : 0.01 min 0.0

End Time : 31.91 min

Plot Offset: 42 mV

Sample #: DSL

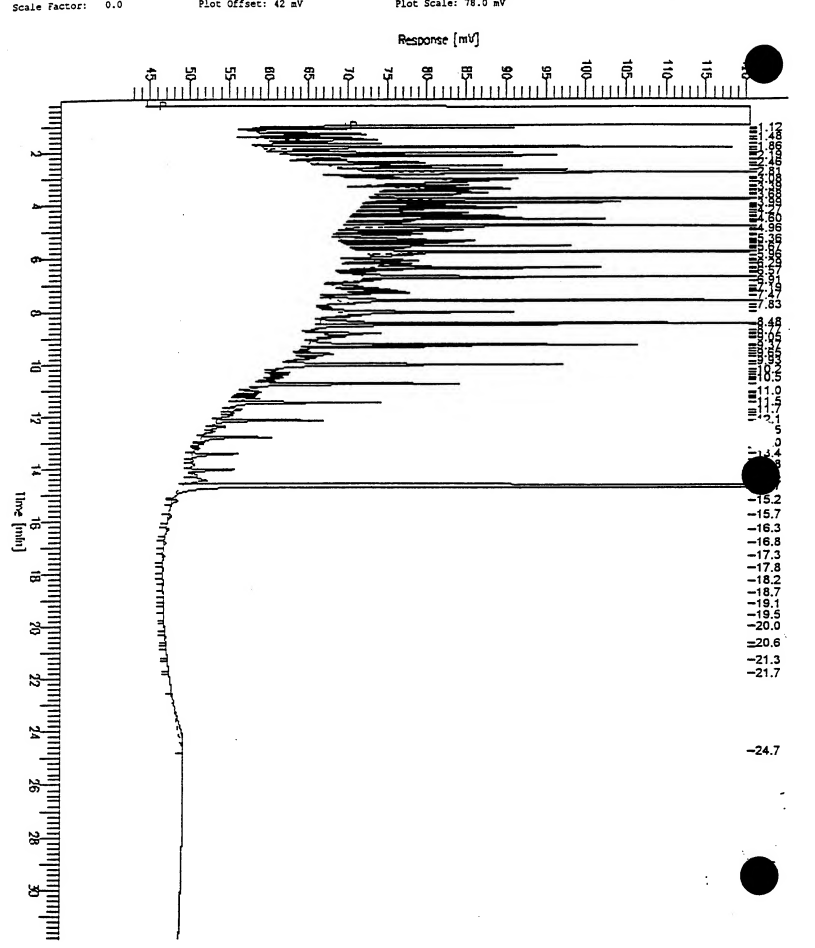
Page 1 of 1

Date: 3/8/96 01:45 PM

Time of Injection: 3/8/96 05:52 AM Low Point : 42.49 mV

High Point : 120.49 mV

Plot Scale: 78.0 mV



GC15 Channel A TEH motor oil

Sample Name: MD 95V5 1596

FileName: C:\GC15\CHB\067B008.RAW >oong/ Sample #: Page 1 of 1 Date: 3/8/96 02:01 PM Time of Injection: 3/7/96 08:23 PM : BTEH.MTH Method End Time : 31.91 min Plot Offset: 42 mV Start Time : 0.01 min Low Point : 42.50 mV High Point : 86.16 mV Factor: 0.0 Plot Scale: 43.7 mV Response [mV] -6.87 -22.4 **≘23.1**



TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M

Project#: 722450.14020

Prep Method: EPA 5030

Location:	Beale	AFB
-----------	-------	-----

Sample # Clie	ent ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124647-001 BE-	MP23-6.5	26407	03/01/96	03/14/96	03/14/96	12%
124647-002 BE-N	4P01-5	26345	03/01/96	03/11/96	03/11/96	12%
124647-003 BE-N	MP43-7.0	26345	03/01/96	03/12/96	03/12/96	13%

Analyte Diln Fac:	Units	124647-001 1	124647-002 1	124647-003 1	
Gasoline	mg/Kg	<1.1	<1.1	<1.1	
Surrogate					
Trifluorotoluene	%REC	93	92	90	
Bromobenzene	%REC	86	81	78	



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M) EPA 5030

Project#: 722450.14020 Prep Method:

Location: Beale AFB

METHOD BLANK

Matrix: Soil Batch#: 26345 Prep Date: 03/11/96 Analysis Date: 03/11/96

Units: mg/Kg Diln Fac: 1

MB Lab ID: QC16891

Analyte	Result	
Gasoline	<1.0	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene Bromobenzene	93 78	52-127 47-112



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Prep Method:

Analysis Method: CA LUFT (EPA 8015M)

EPA 5030

METHOD BLANK

Matrix: Soil Batch#: 26407 Units: mg/Kg Prep Date: Analysis Date:

03/13/96 03/13/96

Diln Fac: 1

MB Lab ID: QC17054

Analyte	Result	
Gasoline	<1.0	
Surrogate	%Rec .	Recovery Limits
Trifluorotoluene Bromobenzene	87 74	52-127 47-112



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method:

EPA 5030

Location: Beale AFB

Soil

LABORATORY CONTROL SAMPLE

Prep Date:

03/11/96

Batch#: 26345 Units: mg/Kg Analysis Date:

03/11/96

Diln Fac: 1

Matrix:

LCS Lab ID: QC16892

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	10.1	10	101	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	106 90	52-127 47-112		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

pike Recovery: 0 out of 1 outside limits



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Soil Batch#: 26407 Units: mg/Kg Prep Date: 03/13/96 Analysis Date:

03/13/96

Diln Fac: 1

LCS Lab ID: QC17055

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	9.1	10	91	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	96 86	52-127 47-112		· · · · · · · · · · · · · · · · · · ·

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits

BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

client: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Method: EPA 5030

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Sample Date: 03/05/96
Lab ID: 124735-013 Received Date: 03/08/96
Matrix: Soil Prep Date: 03/13/96

Batch#: 26407
Units: mg/Kg dry weight
Analysis Date: 03/13/96
Moisture: 9%

Diln Fac: 1

MS Lab ID: QC17057

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Gasoline	10.99	<1.099	10.11	92	75-125
Surrogate	%Rec	Limits			7, 7
Trifluorotoluene Bromobenzene	100 89	52-127 47-112			

CD Lab ID: QC17058

malyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Gasoline	10.99	10	91	75-125	1	<20
Surrogate	%Rec	Limit	s			
Trifluorotoluene Bromobenzene	101 91	52-12 47-11	•	•		

[#] Column to be used to flag recovery and RPD values with an asterisk

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits RPD: 0 out of 1 outside limits



BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Analysis Method: EPA 8020 Prep Method: EPA 5030

Location: Beale AFB

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124647-001	BE-MP23-6.5	26407	03/01/96	03/14/96	03/14/96	12%
124647-002	BE-MP01-5	26345	03/01/96	03/11/96	03/11/96	12%
124647-003	BE-MP43-7.0	26345	03/01/96	03/12/96	03/12/96	13%

Analyte	Units	124647-001	124647-002	124647-003	
Diln Fac:		1	1	1	
Benzene	ug/Kg	<5.7	<5.7	<5.7	
Toluene	ug/Kg	<5.7	<5.7	<5.7	
Ethylbenzene	ug/Kg	<5.7	<5.7	<5.7	
m,p-Xylenes	ug/Kg	<5.7	<5.7	<5.7	
o-Xylene	ug/Kg	<5.7	<5.7	<5.7	
1,2,3-Trimethylbenze	ene ug/Kg	<5.7	<5.7	<5.7	
1,2,4-Trimethylbenze	ene ug/Kg	<5.7	<5.7	<5.7	
1,3,5-Trimethylbenze	ene ug/Kg	<5.7	<5.7	<5.7	
Surrogate					. N. 2-24
Trifluorotoluene	%REC	89	88	87	
Bromobenzene	%REC	80	76	73	

BATCH QC REPORT

Page 1 of 1

BTXE

lient: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

METHOD BLANK

Matrix: Soil Batch#: 26345

Prep Date: 03/11/96
Analysis Date: 03/11/96

Units: ug/Kg Diln Fac: 1

MB Lab ID: QC16891

Analyte	Result	
Benzene	<5.0	
Toluene	<5.0	
Ethylbenzene	<5.0	
m,p-Xylenes	<5.0	
o-Xylene .	<5.0	
1,2,3-Trimethylbenzene	<5.0	
1,2,4-Trimethylbenzene	<5.0	
1,3,5-Trimethylbenzene	<5.0	
Surrogate	%Rec	Recovery Limits

dirogace	*Rec	Recovery Limits
rifluorotoluene	87	43-114
romobenzene	71	45-140



BATCH QC REPORT

Page 1 of 1

			BTX	E		
Project#:	Parsons Engineering S 722450.14020 Beale AFB	Science,	Inc.	Analysis Method: Prep Method:	EPA 8020 EPA 5030	
			METHOD	BLANK		
Matrix: Batch#: Units: Diln Fac:	Soil 26407 ug/Kg 1			Prep Date: Analysis Date:	03/13/96 03/13/96	

MB Lab ID: QC17054

Analyte	Result	
Benzene	<5.0	
Toluene	<5.0	
Ethylbenzene	<5.0	
n,p-Xylenes	<5.0	
o-Xylene	<5.0	•
1,2,3-Trimethylbenzene	<5.0	
1,2,4-Trimethylbenzene	<5.0	
1,3,5-Trimethylbenzene	<5.0	
Surrogate	%Rec	Recovery Limits
Crifluorotoluene	86	43-114
romobenzene	72	45-140



BATCH QC REPORT

Page 1 of 1

BTXE

client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Analysis Method: EPA 8020

Location: Beale AFB

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Soil Batch#:

26345

03/11/96 Prep Date: Analysis Date: 03/11/96

Units: ug/Kg Diln Fac: 1

LCS Lab ID: QC16893

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	106	100	106	80-120
Toluene	109	. 100	109	80-120
Ethylbenzene	105	100	105	80-120
m,p-Xylenes	214	200	107	80-120
o-Xylene	109	100	109	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	89	43-114		
Bromobenzene	74	45-140		

olumn to be used to flag recovery and RPD values with an asterisk lues outside of QC limits

ke Recovery: 0 out of 5 outside limits



BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Soil Batch#: 26407 Units: ug/Kg

Prep Date: 03/14/96 Analysis Date: 03/14/96

Diln Fac: 1

LCS Lab ID: QC17056

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	101	100	101	80-120
Toluene	103	100	103	80-120
Ethylbenzene	99	100	99	80-120
m,p-Xylenes	198	200	99	80-120
o-Xylene	103	100	103	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	85	43-114	 	
Bromobenzene	66	45-140		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 5 outside limits



BATCH QC REPORT

Page 1 of 1

BTXE

client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Analysis Method: EPA 8020

Prep Method:

EPA 5030

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Lab ID: 124736-001

Matrix: Soil Batch#: 26345

Units: ug/Kg dry weight

Diln Fac: 1

Sample Date: 02/26/96 Received Date: 02/29/96

Prep Date: 03/11/96 03/11/96 Analysis Date:

Moisture:

88

MS Lab ID: QC16894

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Benzene	108.7	<5.435	109.8	101	75-125
Toluene	108.7	<5.435	105.4	97	75-125
Ethylbenzene	108.7	<5.435	83.7	77	75-125
m,p-Xylenes	217.4	<5.435	157.6	72 *	75-125
o-Xylene	108.7	<5.435	115.2	106	75-125
Surrogate	%Rec	Limits			
Trifluorotoluene	93	43-114			
romobenzene	69	45-140			

MSD Lab ID: QC16895

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Benzene	108.7	98.91	91	75-125	10	<20
Toluene	108.7	90.22	83	75-125	16	<20
Ethylbenzene	108.7	71.74	66 *	75-125	15	<20
m,p-Xylenes	217.4	132.6	61 *	75-125	17	<20
o-Xylene	108.7	108.7	100	75-125	6	<20
Surrogate	%Rec	Limit	3			
Trifluorotoluene	93	43-114	1			
Bromobenzene	70	45-140	_			

[#] Column to be used to flag recovery and RPD values with an asterisk

Spike Recovery: 3 out of 10 outside limits

^{*} Values outside of QC limits RPD: 0 out of 5 outside limits

PARSONS ENGINEERING SCIENCE, INC.

1301 Marina Village Pkwy, Suite 200 Alameda, California 94501

FAX: (510) 769-9244

Phone: (510) 769-0100

CHAIN OF CUSTODY RECORD

OF \ PAGE

Do MS/MSD sample handbook requirements. FAX COC Upon pecapt to site Manager, er This 4. TMBs: 1,2,3-timethylbenzene, 1,2,4-timethylbenzene, and 1,3,5**according to AFCEE** Note: All analyses and holding times Remarks 9 trimethylbenzene. OTHER INORGANICS **LEVE** Secondary column confirmation required.
 Quantify all carbon ranges: diesel, jet fuel, oll/grease. (ENIP.) 3. Delonized water extraction
DATE TIME REMARKS/COMMENTS: ORGANICS ANALYTES (METHOD) NOTES X NO. OF CONTAINERS RECEIVED BY: (SIGNATURE) Matrix soil soil Soil soil Soil Soil soll **30** soil soll Sol Soil **80** RECEIVED FOR End Depth 40 5.5 5世 Begin Depth 651 5.0 7.0 Project Manager: Todd Wiedemeier (Denver, CO) 3/1/26/1600 2630 Project Name/Location: Beale AFB, UST Site 08 35 Site Manager: Michael Phelps (Alameda, CA) 000/ Time DATE DATE Sampler(s): (Initials and Signature(s)) \mathcal{MBP} $\mathcal{MJLMCMM}$ 3/1/96 3 BE-MP43-7.0 3/1196 3/1/96 Date BELINGUISHED BY: (SIGNATURE) RELINQUISHED BY: (SIGNATURE) Project No.: 722450.14020 GE = EXPAI - S-BE-MP23-65 BE-MPØ1-5 Sample ID

Distribution: Original accompanies shipment; photocopy kept by samplers; copy FAX ed to Parsons ES Site Manager by laboratory upon receipt or samples.

Beale

COOLER RECEIPT CHECKLIST

Curtis	æ	Tompkins, Ltd	
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Login	#: 124647 Date Received: 3/2/96 Number of Coolers:	
Client:		
A.	Preliminary Examination Phase	
	Date Opened: 32 96 By (print): J. Goyett (sign)	
1.	Did cooler come with a shipping slip (airbill, etc.)?	NO
	If YES, enter carrier name and airbill number: FLD Ex 1154090965	
2.	Were custody seals on outside of cooler?	ИО
	How many and where? KRONT Seal date: 31 Seal name:	
3.	Were custody seals unbroken and intact at the date and time of arrival?	
4.	Were custody papers dry and intact when received?	МО
5.	Were custody papers filled out properly (ink, signed, etc.)?	NO
6.	Did you sign the custody papers in the appropriate place?	
7.	Was project identifiable from custody papers?	NO
	If YES, enter project name at the top of this form.	
8.	If required, was sufficient ice used?	NO
	Type of ice: Temperature: 3:2°C	
	, -	
B.	Login Phase	
	Date Logged In: By (print): \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	
1.	Describe type of packing in cooler:	
2.	Did all bottles arrive unbroken?	
3.	Were labels in good condition and complete (ID, date, time, signature, etc.)?	
4.	Did bottle labels agree with custody papers?	
٥.	Were appropriate containers used for the tests indicated?	
6.	Were correct preservatives added to samples?	
7.	Was sufficient amount of sample sent for tests indicated?	
3.	Were bubbles absent in VOA samples? If NO, list sample Ids below	
9.	Was the client contacted concerning this sample delivery?	NO
	If YES, give details below.	
	Who was called? By whom? Date:	
		•
Additio	onal Comments:	•
		
		
Filename:	F:\qc\forms\cooler.wpd Rev. 1 4/95	



Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 9471O, Phone (510) 486-0900

ANALYTICAL REPORT

Prepared for:

Parsons Engineering Science, Inc. 1301 Marina Village Parkway Suite 200 Alameda, CA 94501

Date: 28-MAR-96

Lab Job Number: 124746

Project ID: 722450.14020 Location: Beale AFB

Reviewed by:

Reviewed by:

This package may be reproduced only in its entirety.



Laboratory Number: 124746 Sample Date: 03/06,07/96

Client: Parsons Engineering Science Receipt Date: 03/08/96

Project#: 722450.14020 Location: Beale AFB

CASE NARRATIVE

Curtis & Tompkins, Ltd. received six water samples and one trip blank from the above referenced project on March 8, 1996. All samples were received cold and intact.

TPH/Extractable: No analytical problems were encountered.

TPH/Gasoline: Trifluorotoluene and bromobenzene failed surrogate recovery limits for sample BE-MP14-01 (C&T# 124746-002) due to coelution of sample hydrocarbons with these surrogates. The sample was reanalyzed with similar results, confirming that the failure was due to matrix interferences. The chromatogram for this sample is included in this report. The trip blank was not documented on the chain of custody. The sampling date on the trip blank label was 2/29/96. The analysis was not performed until 3/20/96, which is past the technical holding time.

BTXE & TMBs: The trip blank was not documented on the chain of custody. The sampling date on the trip blank label was 2/29/96. The analysis was not performed until 3/20/96, which is past the technical holding time. As BTXE and gasoline are analyzed concurrently, only one set of matrix spikes are performed. The matrix spikes associated with these samples was analyzed for gasoline. The BTXE laboratory control sample passed acceptance criteria.



MEMORANDUM TO FILE

<u>.</u>	JOB NO. 722450. 14050
	FILE DESIGNATION BRALE AFB UST Site
	DATE 4/4/96 TIME
PHONE CALL FROM Michael Phelps	
PHONE CALL TO	PHONE NO
CONFERENCE WITH	
PLACE	
SUBJECT Trip Blanks, Data Package	25 124746, 124747
The dates on The labels use	I for trip blanks submitted
on The Chain- of- costodies dated	3/7/96 and 3/8/96 were
incorrect. They should have reflected	I The lates on The chain-D-
costody. Therefore, The analyse	s of both trip blanks is
within The analytical method	hold time
	· · · · · · · · · · · · · · · · · · ·
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Chromatograms are included in this report for any samples containing quantifiable concentrations of extractable or purgeable hydrocarbons, along with example chromatograms for each type of fuel.

Data Qualifier Flags

C: The "C" qualifier flag indicates that the target compound reported was confirmed by a second analytical column, but that the concentration quantitated on the second column differed from the reported concentration by more than a factor of two. This often happens when the fuel pattern present in the samples does not resemble gasoline.

DO: The surrogate recovery could not be calculated due to the dilution required for sample analysis.

L: The "L" qualifier flag indicates that the sample chromatogram resembles a fuel pattern, but that the hydrocarbons present are lighter than those seen in a standard of the reported fuel.

H: The "H" qualifier flag indicates that the sample chromatogram resembles a fuel pattern, but that the hydrocarbons present are heavier than those seen in a standard of the reported fuel.

Y: The "Y" qualifier flag is used to indicate that the sample chromatogram does not resemble the standard for the reported fuel range. This flag is often used in combination with the flags "L" and/or "H".

Z: The "Z" qualifier flag indicates that the sample chromatogram does not resemble a fuel pattern, but consists of individual, strongly differentiated peaks.

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1301 Marina Village Pkwy, Sulte 200 Alameda, California 94501 Phone: (510) 769-0100 FAX: (510) 769-9244

THAT F CUSTODY RECORD



Project No.: 722450.14020							ORGANICS	 	INORGANICS	/отнев/
Project Name/Location: Beale AFB, UST Site	UST Site					-	1 / / / / / / / / / / / / / / / / / / /			
Project Manager: Todd Wiedemeier (Denver, CO)	(Denver, CO	~			~O _K	16	96100	///		
Site Manager: Michael Phelps (Alameda, CA)	eda, CA)				MET	200	963 (/	////	
Sampler(s): (Initials and Signature(s))	The Mills	100	The the		TESA	\$100	0208/0			
Ann S			J. J. J. J. J. J. J. J. J. J. J. J. J. J	0. OF C	(JANA Devocoz) or Devocoz) or		Cossos Sant			Note: All analyses and holding times
Sample ID Date	Time	트루	End Depth Matrix		POSEB-HOL	bre X37	AOC & (32)			according to AFCEE handbook requirements. Remarks
BE-MP-2701 3/6/96	1540		water	Ъ	X	7	6 5			TMB's see note 4.
BE-NP14-01 3/6 A6	225	\ \ \ \	water	4	X	X				
3 BE-MPZ6 -01 3/6 A6	1425	l	water	4	又又	X				
BE-NAZA-O1 3/6/96	0530	1	water	8	XX	又				
BE-NP 63-01 3/7/96	1136	1	water	4	X	X				
BE-NP 25-01 3/7 96	C945	(- water	7	X	Z				×
			water							
1 Trio 10-16			water							1.7. BC
49 20 1			water							NON ICH
			water						GEC	ar Selve
			water						TEN PORTO	, ac
			water						D THE	
			water							
RELINQUISHED BY: (SIGNATURE) DATE	E TIME	RECEIVE	RECEIVED BY: (SIGNATURE)	RE)	NOTES:	ES:				
					~ <u>~ ~ ~ ~</u>	econdary Nantify e elonized	 Secondary column confirmation required. Quantify all carbon ranges: diesel, jet fuel, oll/grease Deionized water extraction 	ion required. Iesel, jet fuel, oli/gr		4. TMBs: 1,2,3-trimethylbenzene, 1,2,4- trimethylbenzene, and 1,3,5- trimethylbenzene.
RELINGUISHED BY: (SIGNATURE) DATE	E 138E	RECEIVE	RECEIVED FOR LAB BY: (S	(SIGNATURE)	-	F ,	TIME REMARKS	REMARKS/COMMENTS:		
Marge 17	"HAGHOU		moran.	4	1	20/2	469			
						1				

Pistribution: Original accompanies shipment; photocopy kept by samplers; copy FAXed to Parsons ES Site Manager by laboratory upon receipt of samples.

Berle

COOLER RECEIPT CHECKLIST Curis & Tompkins, Ltd.

Login#: 124746 Date Received: 3/8 Number of Coolers:	
Client: Pusar Project: Beale 653	
A. Preliminary Examination Phase Date Opened: See By (print): (sign) Did cooler come with a shipping slip (airbiil, etc.)?	
Date Opened: By (print): (sign)	
1. Did cooler come with a shipping slip (airbill, etc.)?	
If YES, enter carrier name and airbill number: 7ed 4 115409277	
Were custody seals on outside of cooler?	
How many and where? Seal date: Seal name:	7
Were custody seals unbroken and intact at the date and time of arrival?	1
Were custody papers dry and intact when received?	1
Ware custody papers filled out properly (ink signed etc.)?	
6. Did you sign the custody papers in the appropriate place?	
7. Was project identifiable from custody papers?	\
If YES, enter project name at the top of this form.	\
8. If required, was sufficient ice used?	1
Type of ice: Temperature:	
B. Login Phase 3/	
Date Logged In: 3/8 By (print): 2011 (sign) Les Olle	3
Describe type of packing in cooler:	
2. Did all bottles arrive unbroken? NO	
Were labels in good condition and complete (ID, date, time, signature, etc.)? S NO	
Did bottle labels agree with custody papers? NO	1
5. Were appropriate containers used for the tests indicated?	
6 Were correct preservatives added to samples? NO	- 1
7 Was sufficient amount of sample sent for tests indicated?	1
Were bubbles absent in VOA samples? If NO, list sample Ids below	
9 Was the client contacted concerning this sample delivery?	}
If YES, give details below.	
Who was called? By whom? Date:	. /
Additional Comments:	
Trip blan Louisid	
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CURTIS & TOMPKING 1 TO BEDKE EV	
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C)

LOGIN CHANGE FORM

,					 						
724	Duedate		2/26	7							
11.8 Initials: 74.5	Analysis		7/4/4/T	100		·					
Date/Time: 3/18	Add/Cancel		Add								
	Matrix		H,0								
Client Request: By: M.Phulos Login Review Data Review	Client ID	-	TRIP BLANK					-	-	-	
Client Reque	Previous Lab ID										
Reason for change:	Current Lab ID		t-9/1/61								

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		-	-							
				•						



TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124746-001 BE-MP27-01	26514	03/06/96	03/20/96	03/20/96	
124746-002 BE-MP14-01	26514	03/06/96	03/20/96	03/20/96	
124746-003 BE-MP26-01	26514	03/06/96	03/20/96	03/20/96	
124746-004 BE-MP24-01	26514	03/06/96	03/20/96	03/20/96	

Analyte Diln Fac:	Units	124746-001 1	124746- 1	002	124746-003 1	124746-00 1
Gasoline	ug/L	<50	4000	Y	<50	<50
Surrogate						
Trifluorotoluene	%REC	97	169	*	98	97
Bromobenzene	%REC	95	129	*	97	96

^{*} Values outside of QC limits

Y: Sample exhibits fuel pattern which does not resemble standard



TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124746-005 BE-MP03-01	26514	03/07/96	03/20/96	03/20/96	
124746-006 BE-MP25-01	26514	03/07/96	03/21/96	03/21/96	
124746-007 TRIP BLANK	26514	02/29/96	03/20/96	03/20/96	

Analyte Diln Fac:	Units	124746-005 1	124746-006 1	124746-007 1	
Gasoline	ug/L	<50	<50	<50	
Surrogate					
Trifluorotoluene	%REC	97	98	99	
Bromobenzene	%REC	95	94	94	



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 5030

METHOD BLANK

Matrix: Water Batch#: 26514 Units: ug/L

Diln Fac: 1

Prep Date: 03/20/96 Analysis Date: 03/20/96

MB Lab ID: QC17470

Analyte	Result	
Gasoline	<50	OF THE MARKET TO THE PARTY OF T
Surrogate	%Rec	Recovery Limits
Trifluorotoluene Bromobenzene	99 93	69-120 70-122



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Method: EPA 5030 Location: Beale AFB

LABORATORY CONTROL SAMPLE

Matrix: Water 03/20/96 Prep Date: Batch#: 26514 Analysis Date: 03/20/96 Units: ug/L

Diln Fac: 1

LCS Lab ID: QC17477

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	1979	2000	99	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	104	69-120		
Bromobenzene	107	70-122		

[#] Column to be used to flag recovery and RPD values with an asterisk

pike Recovery: 0 out of 1 outside limits

^{*} Values outside of QC limits



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB

124811-007

Analysis Method: CA LUFT (EPA 8015M) Prep Method:

EPA 5030

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

03/13/96 Sample Date: Received Date: 03/14/96 Prep Date: 03/20/96

Water 26514 Analysis Date: 03/20/96

Units: ug/L Diln Fac: 1

Field ID: ZZZZZZ

Lab ID:

Matrix:

Batch#:

MS Lab ID: QC17472

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Gasoline	2000	<50.00	2004	100	75-125
Surrogate	%Rec	Limits			
Trifluorotoluene Bromobenzene	105 107	69-120 70-122			

MSD Lab ID: QC17473

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Gasoline	2090	2000	105	75-125	5	<20
Surrogate	%Rec	Limits				
Trifluorotoluene Bromobenzene	106 113	69-1: 70-1:			de,	

[#] Column to be used to flag recovery and RPD values with an asterisk

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits RPD: 0 out of 1 outside limits

FileName : G:\GC05\080H018.raw Start Time : 0.00 min Er

Scale Factor: -1

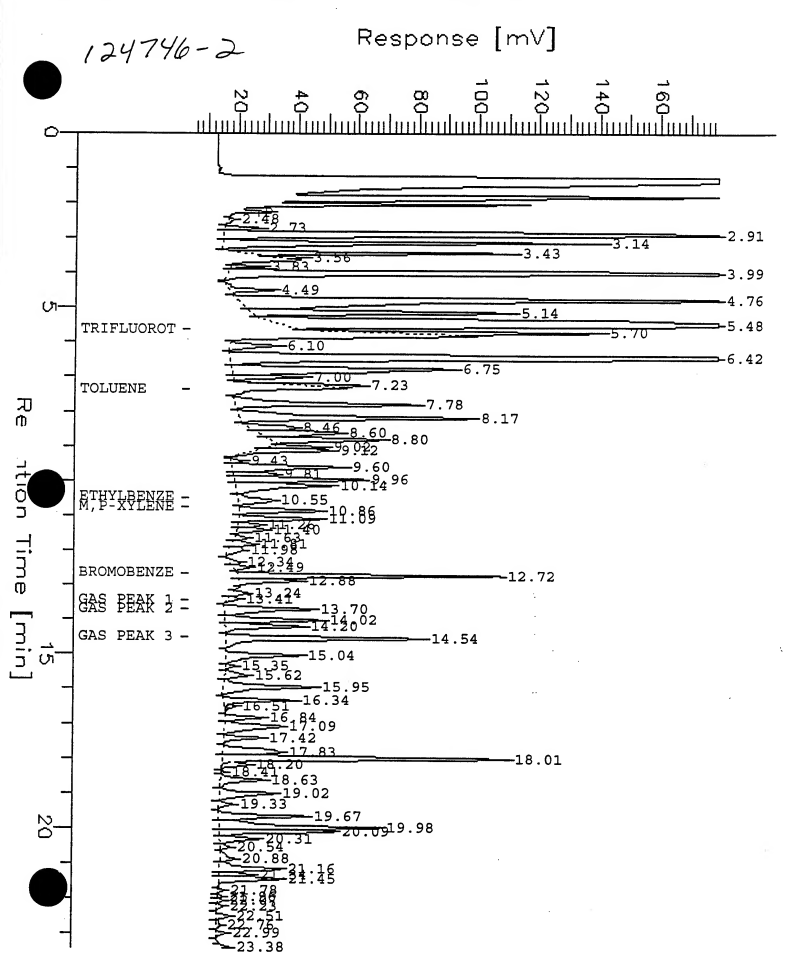
End Time : 23.42 min

Plot Offset: 4 mV

Date: 3/20/96 10:24 PM

Low Point : 4.20 mV Plot Scale: 175 mV

Page 1 of 1 High Point: 179.20 mV



: G:\GC04\135J001.raw FileName Date: 5/15/95 9:53 AM Page 1 of 1 Start Time : 0.00 min End Time : 17.00 min Low Point : 42.92 mV High Point : 217.92 mV Plot Offset: 43 mV Scale Factor: -1 Plot Scale: 175 mV Response [mV] 0.87 1.15 1.35 1.60 1.83 TRIFLUOROT -2.85 -3.02 TOLUENE -3.39 3.91 -4.18 ETHYLBENZE -5.04 -5.20 Retention 5.60 -5.96 BROMOBENZE б.: 6.49 GAS PEAK 1 = 7.14 GAS PEAK 3 -7.70 7.87 8.06 Time . 62 -12.77 14.54 16.10 16.45

IVH2 - GC-04 RTX-1

C+T Gasoline Std.

FileName : G:\GC07\271F031.raw

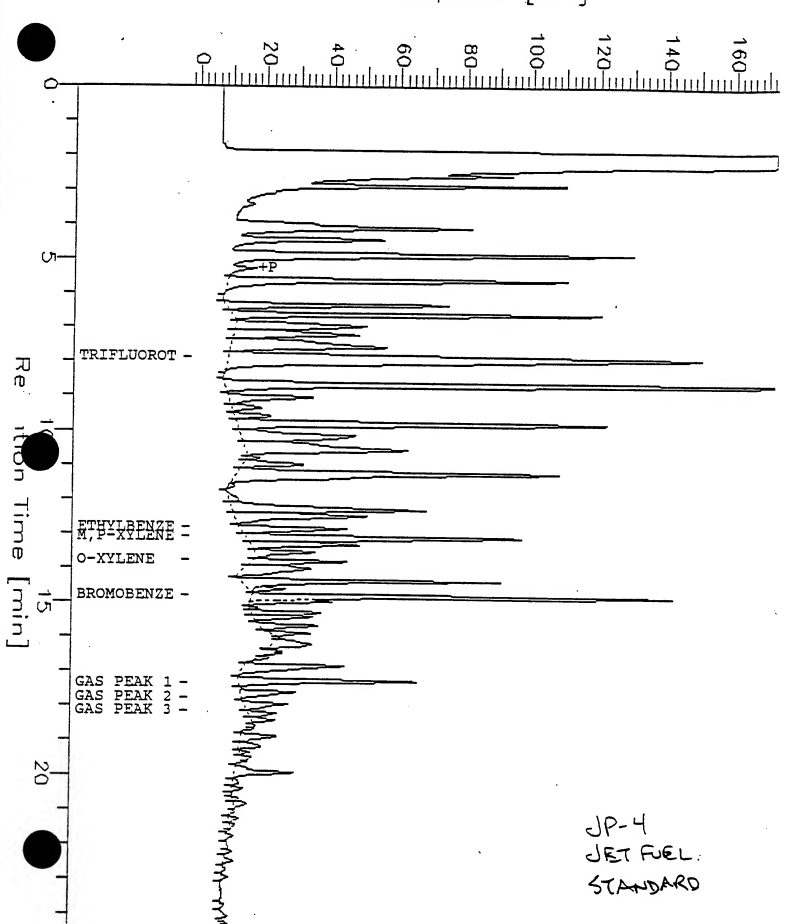
Start Time : 0.00 min Scale Factor: -1

End Time : 24.33 min Plot Offset: -2 mV Date: 9/29/93 9:25 AM Low Point: -2.14 mV

Plot Scale: 175 mV

Page 1 of 1 High Point : 172.86 mV

Response [mV]





BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

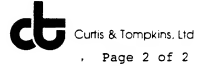
Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124746-001	BE-MP27-01	26514	03/06/96	03/20/96	03/20/96	
124746-002	BE-MP14-01	26514	03/06/96	03/20/96	03/20/96	
124746-003	BE-MP26-01	26514	03/06/96	03/20/96	03/20/96	
124746-004	BE-MP24-01	26514	03/06/96	03/20/96	03/20/96	

	nits	124746-001	124746-002	124746-003	124746-004
Diln Fac:		1	1	1	1
Benzene	ug/L	<0.5	9.1C	<0.5	<0.5
Toluene	ug/L	<0.5	30 C	<0.5	<0.5
Ethylbenzene	ug/L	<0.5	10	<0.5	<0.5
m,p-Xylenes	ug/L	<0.5	19	<0.5	<0.5
o-Xylene	ug/L	<0.5	3.8	<0.5	<0.5
1,2,3-Trimethylbenzene	ug/L	<0.5	24	<0.5	<0.5
1,2,4-Trimethylbenzene	ug/L	<0.5	24	<0.5	<0.5
1,3,5-Trimethylbenzene	ug/L	<0.5	36	<0.5	<0.5
Surrogate					
Trifluorotoluene	%REC	96	91	97	95
Bromobenzene	%REC	93	89	95	94

C: Presence of this compound confirmed by second column, however, the confirmation concentration differed from the reported result by more than a factor of two



BTXE

lient: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124746-005 BE-MP03-01	26514	03/07/96	03/20/96	03/20/96	
124746-006 BE-MP25-01	26514	03/07/96	03/21/96	03/21/96	
124746-007 TRIP BLANK	26514	02/29/96	03/20/96	03/20/96	

Analyte Diln Fac:	Units	124746-005 1	124746-006 1	124746-007 1	
Benzene	ug/L	<0.5	<0.5	<0.5	
Toluene	ug/L	<0.5	<0.5	<0.5	
Ethylbenzene	ug/L	<0.5	<0.5	<0.5	
m,p-Xylenes	ug/L	<0.5	<0.5	<0.5	
o-Xylene	ug/L	<0.5	<0.5	<0.5	
1,2,3-Trimethylbenzene	e ug/L	<0.5	<0.5	<0.5	
2,4-Trimethylbenzene	e ug/L	<0.5	<0.5	<0.5	
3,5-Trimethylbenzene	e ug/L	<0.5	<0.5	<0.5	
Surrogate					
Trifluorotoluene	%REC	95	95	97	
Bromobenzene	%REC	93	94	93	



Lab #: 124746

BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

METHOD BLANK

Matrix: Water Prep Date: 03/20/96 Batch#: 26514 Analysis Date: 03/20/96

Units: ug/L Diln Fac: 1

MB Lab ID: QC17470

Analyte	Result	
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene	99	58-130
Bromobenzene	94	62-131

BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26514

Prep Date: 03/20/96 Analysis Date: 03/20/96

Units: ug/L Diln Fac: 1

LCS Lab ID: QC17471

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	18	20	90	80-120
Toluene	19	20	95	80-120
Ethylbenzene	18	. 20	90	80-120
m,p-Xylenes	36	40	90	80-120
o-Xylene	18	20	90	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	100	58-130		· · · · · · · · · · · · · · · · · · ·
Bromobenzene	95	62-131		

Column to be used to flag recovery and RPD values with an asterisk values outside of QC limits

ke Recovery: 0 out of 5 outside limits



TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M

Project#: 722450.14020

Prep Method: EPA 3520

Location: Beale AFB

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124746-001 BE-MP27-01	26390	03/06/96	03/12/96	03/14/96	
124746-002 BE-MP14-01	26390	03/06/96	03/12/96	03/14/96	
124746-003 BE-MP26-01	26390	03/06/96	03/12/96	03/14/96	
124746-004 BE-MP24-01	26390	03/06/96	03/12/96	03/14/96	

Analyte Diln Fac:	Units	124746-001 1	46-001 124746-002 1 1		124746-003 1	124746-004 1
JP5	ug/L	<50	1900	YL	<50	<50
Diesel Range	ug/L	<50	2200	YL	<50	<50
Motor Oil Range	ug/L	<300	310	YL	<300	<300
Surrogate						
Hexacosane	%REC	70	70		75	72

Y: Sample exhibits fuel pattern which does not resemble standard

L: Lighter hydrocarbons than indicated standard



Page 2 of 2

TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 3520

Location: Beale AFB

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124746-005	BE-MP03-01	26390	03/07/96	03/12/96	03/13/96	
124746-006	BE-MP25-01	26390	03/07/96	03/12/96	03/13/96	

Analyte Diln Fac:	Units	124746-005 1	124746-006 1	
JP5	ug/L	<50 ·	<50	
Diesel Range	ug/L	<50	<50	
Motor Oil Range	ug/L	<300	<300	
Surrogate				
lexacosane	%REC	71	67	

Lighter hydrocarbons than indicated standard



BATCH QC REPORT

Page 1 of '

TEH-Tot Ext Hydrocarbons

Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 3520

METHOD BLANK

Matrix: Water Batch#: 26390 Units: ug/L

Prep Date: 03/12/96 Analysis Date: 03/14/96

Diln Fac: 1

1.17

MB Lab ID: QC16985

Analyte	Result	
JP5	<50	
Diesel Range	<50	
Motor Oil Range	<300	
Surrogate	*Rec	Recovery Limits
Hexacosane	69	60-140

BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 3520

Location: Beale AFB

BLANK SPIKE/BLANK SPIKE DUPLICATE

03/12/96

Matrix: Water Batch#: 26390 Units: ug/L

Diln Fac: 1

Prep Date: 0
Analysis Date: 0

03/13/96

BS Lab ID: QC16986

Analyte	Spike Added	BS	%Rec #	Limits
Diesel Range	2475 21	43	. 87	60-140
Surrogate	%Rec :	Limits		
Hexacosane	65	60-140		

BSD Lab ID: QC16987

Analyte	Spike Added	BSD	%Rec #	Limits	RPD #	Limit
iesel Range	2475	2278	92	60-140	6	<35
Surrogate	%Rec	Limit	s			
Hexacosane	68	60-14	0			

[#] Column to be used to flag recovery and RPD values with an asterisk

RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits

TEH Unromatogram - GU II UN B

Sample Name: 124746-002,500:2.5 FileName : g:\gc11\chb\073B040.raw

: 11B32.ins

Method Start Time : 0.00 min Scale Factor: -1

End Time : 31.92 min Plot Offset: 35 mV

Sample #: 26390

Date: 3/14/96 10:15 AM

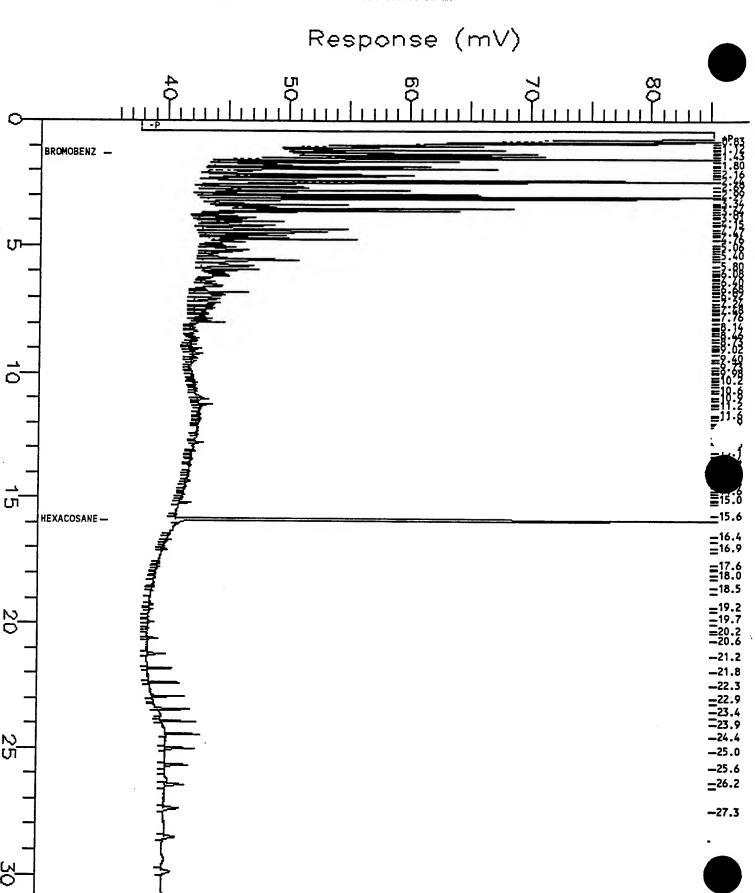
Time of Injection: 3/14/96 09:42 AM

Low Point: 35.21 mV

High Point: 85.21 mV

Page 1 of 1

-31.5



TEH Chromatogram - GC 11 Ch B

Sample Name : DIESEL 500MG/L

FileName : g:\gc11\chb\073B014.raw

Method : 11B20.ins

Start Time : 0.00 min 'le Factor: -1

End Time : 20.00 min

Plot Offset: 37 mV

Sample #: 96WS1948

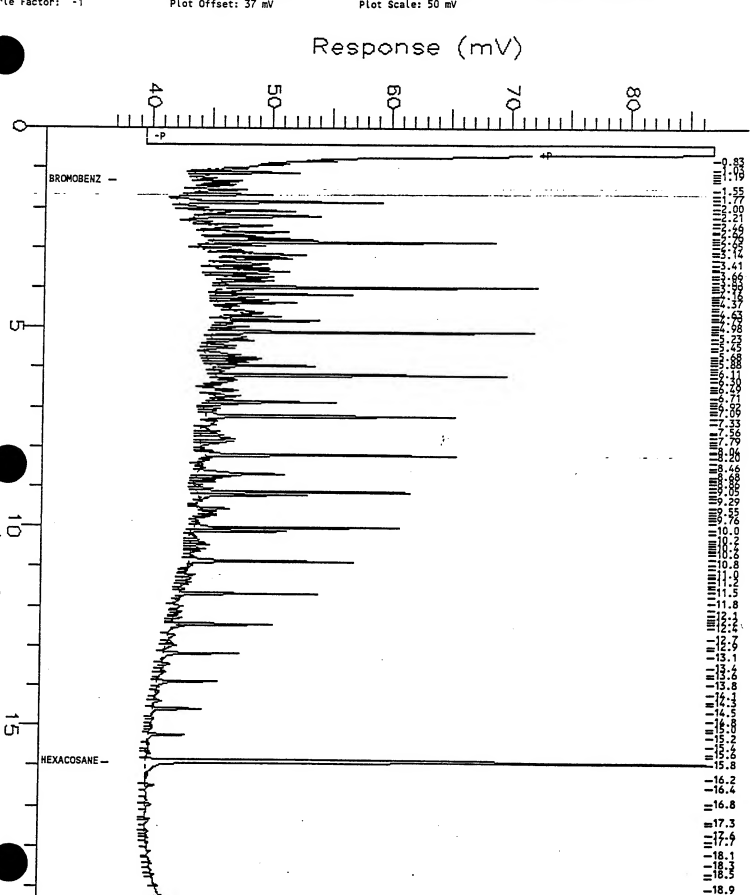
Date: 3/13/96 04:45 PM

Time of Injection: 3/13/96 04:24 PM

Low Point : 36.92 mV

High Point: 86.92 mV

Page 1 of 1



TEH Chromatogram - GC 11 Ch B

Sample Name : JP-5 250MG/L

: g:\gc11\chb\073B015.raw : 11B20.ins FileName

Method

Start Time : 0.00 min Scale Factor: -1

End Time : 20.00 min Plot Offset: 35 mV

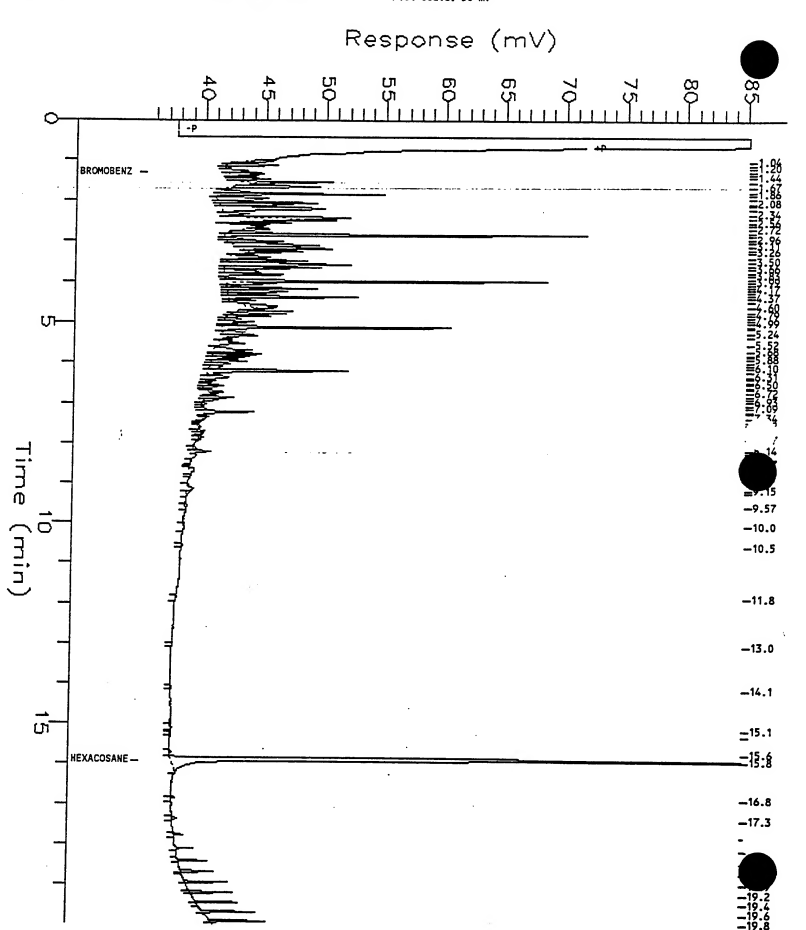
Sample #: 96WS2068 Date : 3/13/96 05:15 PM

Time of Injection: 3/13/96 04:55 PM

Low Point : 35.11 mV

High Point : 85.11 mV

Page 1 of 1



TEH Chromatogram - GC 11 Ch B

Sample Name : MOTOR OIL 500MG/L

FileName : g:\gc11\chb\073B016.raw

Method : 11B32.ins Start Time : 0.00 min Thale Factor: -1

End Time : 31.92 min

Plot Offset: 34 mV

Sample #: 95WS1596

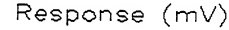
Date: 3/13/96 05:58 PM

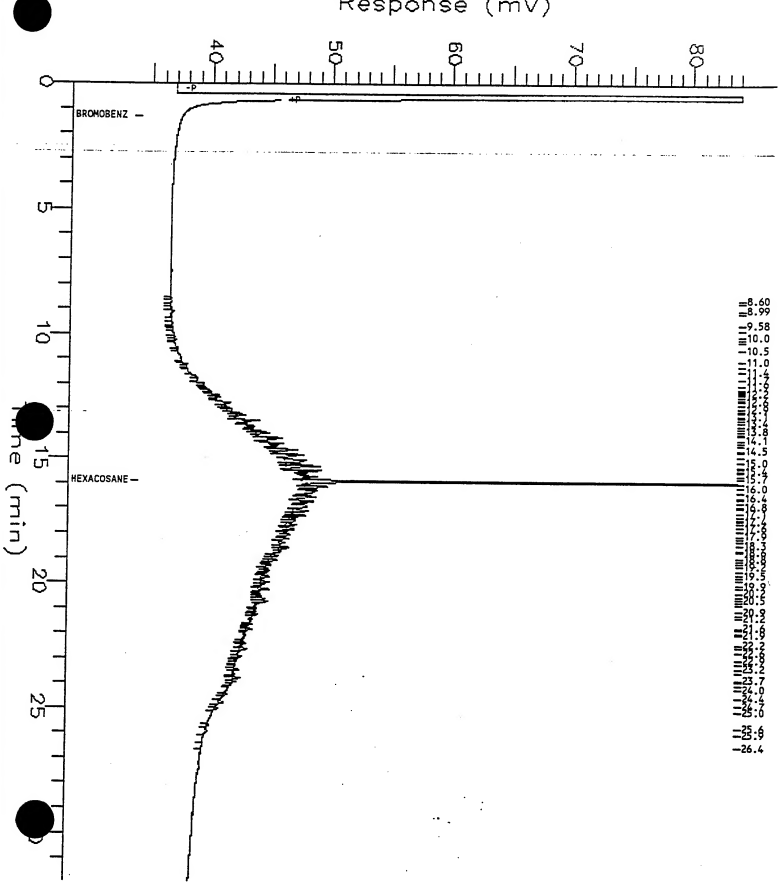
Time of Injection: 3/13/96 05:25 PM

Low Point : 34.04 mV

High Point : 84.04 mV

Page 1 of 1







Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 9471O, Phone (510) 486-0900

ANALYTICAL REPORT

Prepared for:

Parsons Engineering Science, Inc. 1301 Marina Village Parkway Suite 200 Alameda, CA 94501

Date: 28-MAR-96

Lab Job Number: 124747

er kan da Ministra

Project ID: 722450.14020 Location: Beale AFB

Reviewed by: June K

Reviewed by:

This package may be reproduced only in its entirety.



Laboratory Number: Sample Date: 03/07,08/96 124747 Receipt Date: 03/08/96

Client: Parsons Engineering Science

Project#: 722450.14020 Location: Beale AFB

CASE NARRATIVE

Curtis & Tompkins, Ltd. received six water samples and one trip blank from the above referenced project on March 8, 1996. All samples were received cold and intact.

TPH/Extractable: No analytical problems were encountered.

TPH/Gasoline: The trip blank was not documented on the chain of custody. The sampling date on the trip blank label was 2/29/96. Analysis of this trip blank was not completed until 3/22/96, which is past the technical holding time. As BTXE and gasoline are analyzed concurrently, only one set of matrix spikes are performed. The matrix spikes associated with these samples were analyzed for BTXE. The gasoline laboratory control samples passed acceptance criteria.

BTXE & TMBs: The trip blank was not documented on the chain of custody. The sampling date on the trip blank label was 2/29/96. The analysis was not performed until 3/22/96, which is past the technical holding time.



MEMORANDUM TO FILE

	JOB NO. 722450. 14050
	FILE DESIGNATION BRALE AFB UST Site
	DATE 4/4/96 TIME
·	
PHONE CALL FROM Michael Phelps	PHONE NO
PHONE CALL TO	PHONE NO
CONFERENCE WITH	•
	` .
PLACE	
SUBJECT Trip Blanks, Data Package	15 124746, 124747
The dates on The labels use	I for trip blanks submitted
on The Chain- of- custodies dated incorrect. They should have reflected	O The dates on The Challe of
the Tourse Tourse	ALD tight
custody. Therefore, The analyse	5 0/ 601h 1rip blanks 15
within The analytical method	hold time.
	·
	•
3	
	201 2 321

PARSONS ENGINEERING SCIENCE, INC.

1301 Marina Village Pkwy, Suite 200

Phone: (510) 769-0100 FAX: (510) 769-9244

Alameda, California 94501

LECORD LECORD RECORD

OF

PAGE 1

handbook requirements. 4. TMBs: 1,2,3-trimethylbenzene, 1,2,4 trimethylbenzene, and 1,3,5**according to AFCEE** and holding times Note: All analyses Remarks MEND RECEIVED bimethylbenzene. OTHER, Š NORGANICS Secondary column confirmation required.
 Quantity all carbon ranges: diesel, jet fuel, oll/grease.
 Delonized water extraction. REMARKS/COMMENTS: 1 81001 804 (0 1 08/0 £02) ORGANICS \$7.5% and TMBs (5030/8020) saw TMB 1,1 71ME 6:35 BION BBS (MRTOBIOEOR) BANDORDE HATT ANALYTES (METHOD) RECEIVED FOR LAB BY: (SIGNATURE) * NO. OF CONTAINERS 4 RECEIVED BY: (SIGNATURE) D. Moore Water water Water water water water water water water water water water Water Matrix End Depth ١ j Begin Depth l Project Manager: Todd Wiedemeier (Denver, CO) TIME 0220 1430 Project Name/Location: Beale AFB, UST Site 3/4/96 1500 1345 1320 Site Manager: Michael Phelps (Alameda, CA) Time DATE 8/7/96 3/8/26 3/8/96 16/8/2 3/8/8/6 Sampler(s): (Initials and Signature(s) Mrs track H. Fricolum Date RELINQUISHED BY: (SIGNATURE) INQUISHED BY: (SIGNATURE) Project No.: 722450.14020 BE-MW19-01 BE-MUDZ-01 1000 X BE - MW/81-01 BE-MPZG-0/ BE-MW18-01 BE-MP21-01 とっていい Sample ID

Distribution: Original accompanies shipment; photocopy kept by samplers; copy FAXed to Parsons ES Site Manager by laboratory upon receipt of samples.

Berle

COOLER RECEIPT CHECKLIST

Curis & Tompkins, Ltd.

	124747 3/8	l	•
Login#			
Client:			
0		٠	
A.	Preliminary Examination Phase	·• O	0 /
	Date Opened: 3(9) By (print): (sign) (view)	<u>ulk</u>	
1.	Did cooler come with a shipping slip (airbill, etc.)?	<u></u> YEŚ	NO
	If YES, enter carrier name and airbiil number:		<u> </u>
2.	Were custody seals on outside of cooler?		(NO)
_,	How many and where? Seal date: Seal name:		
3.	Were custody seals unbroken and intact at the date and time of arrival?	. YES	NO W
4.	Were custody papers dry and intact when received?	. (YES	NO
5.	Were custody papers filled out properly (ink, signed, etc.)?		NO
б.	Did you sign the custody papers in the appropriate place?	. (3)	NO
7.	Was project identifiable from custody papers?		NO
•	If YES, enter project name at the top of this form.	2	
3.	If required was sufficient ice used?	. E	NO
•	Type of ice: Sylve Temperature: 5,0°C		
В.	Login Phase	\	
-	Date Logged In: 3/4 By (print): delle : (sign) demou	<u> </u>	<u> </u>
1.	Describe type of packing in cooler:		
2.	Did all bottles arrive unbroken?		МО
3.	Were labels in good condition and complete (ID, date, time, signature, etc.)?		ИС
4.	Did bottle labels agree with custody papers?		NO
5.	Were appropriate containers used for the tests indicated?		МО
ó.	Were correct preservatives added to samples?		NO
7.	Was sufficient amount of sample sent for tests indicated?		NO
3.	Were bubbles absent in VOA samples? If NO, list sample Ids below	. YES	NO
9.	Was the client contacted concerning this sample delivery?	. YES	NO
	If YES, give details below.		
	Who was called? By whom? Date	: <u></u>	
Additio	nal Comments:		•
	Tripbluk not un C-000		
	Tripblenk not un (-0-C		
Filename:	F:\qc\forms\cooler.wpd Rev	. 1 4 95	

5		Duedate	2/2	772				•			
LOGIN CHANGE FORM	3/18/56Initials:	Analysis	SWII -	NAC 1VH/15/MTS							
GIN C	Date/Time: 3	Add/Cancol		Add							
		Matrix		HZD							
LTD. BERKELEY	Request: By: M. Phelos Review Data Review	Cllent II)		TRIP BLANK							
	Client Request	Provious Lab ID									
SURTIS & TOMPKINS,	Reason for change:	Current Lab ID	1 1/1-/101	1-12/10							



TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Location: Beale AFB

Prep Method: EPA 3520

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124747-001 BE-MP20-01	26390	03/08/96	03/12/96	03/14/96	
124747-002 BE-MW02-01	26390	03/08/96	03/12/96	03/14/96	•
124747-003 BE-MP21-01	26390	03/08/96	03/12/96	03/14/96	
124747-004 BE-MW01-01	26390	03/08/96	03/12/96	03/14/96	

Analyte Diln Fac:	Units	124747-001 1	124747-002 1	124747-003 1	124747 - 004 1
JP5	ug/L	<50	<50	<50	<50
Diesel Range	ug/L	<50	<50	<50	<50
Motor Oil Range	ug/L	<300	<300	<300	<300
Surrogate					
xacosane	%REC	69	76	76	72



TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M

Project#: 722450.14020

Prep Method: EPA 3520

Location: Beale AFB

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124747-005 BE-MW18-01	26390	03/07/96	03/12/96	03/14/96	

Analyte Diln Fac:	Units	124747-005 1	
JP5	ug/L	<50	
Diesel Range	ug/L	<50	
Motor Oil Range	ug/L	<300	
Surrogate			
Hexacosane	%REC	73	



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Metho

Prep Method: EPA 3520

Location: Beale AFB

METHOD BLANK

Matrix: Water

Prep Date: 03/12/96 Analysis Date: 03/14/96

Batch#: 26390 Units: ug/L Diln Fac: 1

MB Lab ID: QC16985

Analyte	Result	
JP5	<50	
Diesel Range	<50	
Motor Oil Range	<300	
Surrogate	%Rec	Recovery Limits
Hexacosane	69	60-140



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method:

EPA 3520

BLANK SPIKE/BLANK SPIKE DUPLICATE

Matrix: Water Batch#: 26390 Units: ug/L

Diln Fac: 1

Prep Date: Analysis Date: 03/12/96

03/13/96

BS Lab ID: QC16986

Analyte	Spike Added	BS	%Rec #	Limits
Diesel Range	2475	2143	87	60-140
Surrogate	%Rec .	Limits		
Hexacosane	65	60-140		

BSD Lab ID: QC16987

Analyte	Spike Added	BSD	%Rec #	Limits	RPD #	Limi	t
Diesel Range	2475	2278	92	60-140	6	<35	
Surrogate	%Rec	Limit	s				
Hexacosane	68	60-14	10				

[#] Column to be used to flag recovery and RPD values with an asterisk

RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits



Page 1 of 2

TVH-Total Volatile Hydrocarbons

client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124747-001 BE-MP20-01	26544	03/08/96	03/22/96	03/22/96	······································
124747-002 BE-MW02-01	26544	03/08/96	03/22/96	03/22/96	
124747-003 BE-MP21-01	26544	03/08/96	03/22/96	03/22/96	
124747-004 BE-MW01-01	26544	03/08/96	03/22/96	03/22/96	

Analyte Diln Fac:	Units	124747-001 1	124747-002 1	124747-003 1	124747-004
Gasoline	ug/L	<50	<50	<50	<50
Surrogate				44	
Prifluorotoluene	%REC	100	101	100	100
romobenzene	%REC	94	95	95	95



TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: EPA 5030

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124747-005	BE-MW18-01	26544	03/07/96	03/22/96	03/22/96	· · · · · · · · · · · · · · · · · · ·
124747-006	BE-MW19-01	26544	03/07/96	03/22/96	03/22/96	
124747-007	TRIP BLANK	26573	02/29/96	03/22/96	03/22/96	

Analyte Diln Fac:	Units	124747-005 1	124747-006	124747-007 1	
Gasoline	ug/L	<50	<50	<50	
Surrogate					
Trifluorotoluene	%REC	101	102	99	
Bromobenzene	%REC	95	99	95	



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 5030

Project#: 722450.14020 Location: Beale AFB

METHOD BLANK

Matrix: Water Batch#: 26544

Prep Date: 03/21/96 Analysis Date: 03/21/96

Units: ug/L Diln Fac: 1

MB Lab ID: QC17561

Analyte	Result	
Gasoline	<50	
Surrogate	%Rec .	Recovery Limits
Trifluorotoluene Bromobenzene	98 89	69-120 70-122



BATCH QC REPORT

Page 1 of '

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: EPA 5030

METHOD BLANK

Matrix: Water Batch#: 26573 Units: ug/L Diln Fac: 1 Prep Date: 03/22/96 Analysis Date: 03/22/96

MB Lab ID: QC17665

Analyte	Result	
Gasoline	<50	The state of the s
Surrogate	%Rec	Recovery Limits
Trifluorotoluene Bromobenzene	98 91	69-120 70-122



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client:

Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26544 Units: ug/L

Diln Fac: 1

Prep Date:

03/21/96

Analysis Date: 03/21/96

LCS Lab ID: QC17562

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	2008	2000	100	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	104 111	69-120 70-122		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits pike Recovery: 0 out of 1 outside limits



BATCH QC REPORT

Page 1 of '

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26573 Units: ug/L Diln Fac: 1

Prep Date: 03/22/96 Analysis Date: 03/22/96

LCS Lab ID: QC17666

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	2008	2006	100	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	106 112	69-120 70-122	-	

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits



BTXE

lient: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124747-001 BE-MP20-01	26544	03/08/96	03/22/96	03/22/96	
124747-002 BE-MW02-01	26544	03/08/96	03/22/96	03/22/96	
124747-003 BE-MP21-01	26544	03/08/96	03/22/96	03/22/96	
124747-004 BE-MW01-01	26544	03/08/96	03/22/96	03/22/96	

Analyte t Diln Fac:	Jnits	124747-001 1	124747-002 1	124747-003 1	124747-004 1
Benzene	ug/L	<0.5	<0.5	<0.5	<0.5
Toluene	ug/L	<0.5	<0.5	<0.5	<0.5
Ethylbenzene	ug/L	<0.5	<0.5	<0.5	<0.5
m,p-Xylenes	ug/L	<0.5	<0.5	<0.5	<0.5
o-Xylene	ug/L	<0.5	<0.5	<0.5	<0.5
2,3-Trimethylbenzene	ug/L	<0.5	<0.5	<0.5	<0.5
2,4-Trimethylbenzene		<0.5	<0.5	<0.5	<0.5
1,3,5-Trimethylbenzene	ug/L	<0.5	<0.5	<0.5	<0.5
Surrogate					
Trifluorotoluene	%REC	99	97	98	97
Bromobenzene	%REC	94	95	95	95



BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124747-005 BE-MW18-01	26544	03/07/96	03/22/96	03/22/96	
124747-006 BE-MW19-01	26544	03/07/96	03/22/96	03/22/96	
124747-007 TRIP BLANK	26573	02/29/96	03/22/96	03/22/96	

Analyte U	nits	124747-005 1	124747-006 1	124747-007 1	
Benzene	ug/L	<0.5	<0.5	<0.5	
Toluene	ug/L	<0.5	0.6	<0.5	
Ethylbenzene	ug/L	<0.5	<0.5	<0.5	
m,p-Xylenes	ug/L	<0.5	<0.5	<0.5	
o-Xylene	ug/L	<0.5	<0.5	<0.5	
1,2,3-Trimethylbenzene	ug/L	<0.5	<0.5	<0.5	
1,2,4-Trimethylbenzene	ug/L	<0.5	<0.5	<0.5	
1,3,5-Trimethylbenzene	ug/L	<0.5	<0.5	<0.5	
Surrogate					
Trifluorotoluene	%REC	98	98	100	
Bromobenzene	%REC	96	97	91	

BATCH QC REPORT

Page 1 of 1

BTXE

ient: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

roject#: 722450.14020 Location: Beale AFB

Prep Method:

EPA 5030

METHOD BLANK

Matrix:

Diln Fac: 1

Water

Prep Date:

03/21/96

Batch#: 26544 Units: ug/L

Analysis Date: 03/21/96

MB Lab ID: QC17561

Analyte	Result	
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	

Surrogate	%Rec	Recovery Limits
Trifluorotoluene	96	58-130
pmobenzene	91	62-131



BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

METHOD BLANK

Matrix: Water Batch#: 26573

Units: ug/L Diln Fac: 1 Prep Date: 03/22/96 Analysis Date: 03/22/96

MB Lab ID: QC17665

Analyte	Result	
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	

Surrogate	%Rec	Recovery Limits
Trifluorotoluene	80	58-130
Bromobenzene	88	62-131



BATCH QC REPORT

Page 1 of 1

BTXE

lient: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26544 Units: ug/L Diln Fac: 1

Prep Date: 03/21/96 Analysis Date: 03/21/96

LCS Lab ID: QC17563

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	18.5	20	93	80-120
Toluene	18.5	20	93	80-120
Ethylbenzene	18.1	20	91	80-120
m,p-Xylenes	36.5	40	91	80-120
o-Xylene	18.4	20	92	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	97	58-130		
Bromobenzene	91	62-131		

Column to be used to flag recovery and RPD values with an asterisk lues outside of QC limits

e Recovery: 0 out of 5 outside limits



BATCH QC REPORT

Page 1 of '

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26573 Units: ug/L

Diln Fac: 1

Prep Date: 03/22/96 Analysis Date: 03/22/96

LCS Lab ID: QC17667

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	18.5	20	93	80-120
Toluene	18.4	20	92	80-120
Ethylbenzene	18	. 20	90	80-120
m,p-Xylenes	36.6	40	92	80-120
o-Xylene	18.5	20	93	80-120
Surrogate	%Rec	Limits	*	
Trifluorotoluene	100	58-130		
Bromobenzene	90	62-131		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 5 outside limits

BATCH QC REPORT

Page 1 of 1

BTXE

lient: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Lab ID: 124881-001

Matrix: Water Batch#: 26544 Units: ug/L Diln Fac: 1

Sample Date: 03/19/96 Received Date: 03/19/96 Prep Date: 03/21/96

Analysis Date: 03/21/96

MS Lab ID: QC17564

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Benzene	20	<0.5000	18.6	93	75-125
Toluene	20	<0.5000	18.9	95	75-125
Ethylbenzene	20	<0.5000	18.4	92	75-125
m,p-Xylenes	40	<0.5000	37	93	75-125
o-Xylene	20	<0.5000	18.8	94	75-125
Surrogate	%Rec	Limits			
Trifluorotoluene	96	58-130		·	· · · · · · · · · · · · · · · · · · ·
Bromobenzene	90	62-131			

MSD Lab ID: QC17565

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Benzene	20	19.2	96	75-125	3	<20
Toluene	20	19.6	98	75-125	4	<20
Ethylbenzene	20	19.2	96	75-125	4	<20
m,p-Xylenes	40	38.3	96	75-125	4	<20
o-Xylene	20	19.6	98	75-125	4	<20
Surrogate	%Rec	Limit	s	·		
Trifluorotoluene	97	58-13	0			
Bromobenzene	94	62-13	1			

[#] Column to be used to flag recovery and RPD values with an asterisk

Spike Recovery: 0 out of 10 outside limits

^{*} Values outside of QC limits RPD: 0 out of 5 outside limits



BATCH QC REPORT

Page 1 of

Client: Parsons Engineering Science, Inc. Analysis Method: EPA 8020
Project#: 722450.14020 Prep Method: EPA 5030
Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Sample Date: 03/19/96 Lab ID: 124900-009 Received Date: 03/20/96 Matrix: Water Prep Date: 03/22/96 Batch#: 26573 Analysis Date: 03/22/96 Units: ug/L

Diln Fac: 1

MS Lab ID: QC17668

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Benzene Toluene	20	<0.5000	19.1	96	75-125
Ethylbenzene	20 20	<0.5000 <0.5000	19.1 19.1	96 96	75-125 75-125
m,p-Xylenes o-Xylene	40 20	<0.5000 <0.5000	38.1 19.4	95 97	75-125 75-125
Surrogate	%Rec	Limits			
Trifluorotoluene Bromobenzene	101 91	58-130 62-131			

MSD Lab ID: QC17669

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Benzene	20	19.8	99	75-125	4	<20
Toluene	20	19.7	99	75-125	3	<20
Ethylbenzene	20	19.5	98	75-125	2	<20
m,p-Xylenes	40	39	98	75-125	2	<20
o-Xylene	20	19.8	99	75-125	2	<20
Surrogate	%Rec	Limit	s			
Trifluorotoluene	101	58-130)			- <u></u>
Bromobenzene	94	62-13:	1			

[#] Column to be used to flag recovery and RPD values with an asterisk

RPD: 0 out of 5 outside limits

Spike Recovery: 0 out of 10 outside limits

^{*} Values outside of QC limits



Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 94710, Phone (510) 486-0900

ANALYTICAL REPORT

Prepared for:

Parsons Engineering Science, Inc. 1301 Marina Village Parkway Suite 200 Alameda, CA 94501

Date: 28-MAR-96

Lab Job Number: 124796

Project ID: 722450.14020 Location: Beale AFB

Reviewed by:

Reviewed by:

This package may be reproduced only in its entirety.



Laboratory Number: 124796

Client: Parsons Engineering Science Project#: 722450.14020

Location: Beale AFB

Sample Date: 03/11,12/96 Receipt Date: 03/13/96

CASE NARRATIVE

Curtis & Tompkins received six water samples from the above referenced project on March 13, 1996. The chain of custody incorrectly indicated that all samples were soil. No amber liter was received for sample BE-MP15-01. The project manager was notified of these discrepancies on March 13, 1996.

TPH/Extractable: The sample chromatogram is included in this report for sample BE-MP22-01 (C&T# 124796-004), along with example chromatograms for JP-5 and diesel. No analytical problems were encountered.

TPH/Gasoline: Sample chromatograms are included for any sample containing quantifiable concentrations of purgeable hydrocarbons, along with an example gasoline standard. The patterns exhibited in these sample chromatograms do not resemble gasoline, but are similar to JP-4.

BTXE (EPA 8020): The "C" qualifier flag indicates that the target compound reported was confirmed by a second analytical column, but that the concentration quantitated on the second column differed from the reported concentration by more than a factor of two. This often happens when the fuel pattern present in the samples does not resemble gasoline.



TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 3520

Location: Beale AFB

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124796-002 124796-004		26473	•	–	03/21/96	
124/96-004	BE-MP22-U1	26473	03/12/96	03/18/96	03/21/96	

Analyte Diln Fac:	Units	124796-002 1	124796- 1	-004	
JP5	ug/L	<50	340	Y	
Diesel Range	ug/L	<50	320	Y	
Motor Oil Range	ug/L	<300	<300		
Surrogate					
Hexacosane	%REC	103	95		

Sample exhibits fuel pattern which does not resemble standard



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: EPA 3520

METHOD BLANK

 Matrix:
 Water
 Prep Date:
 03/18/96

 Batch#:
 26473
 Analysis Date:
 03/21/96

Units: ug/L Diln Fac: 1

MB Lab ID: QC17319

Result	
<50	
<50	•
<300	
%Rec	Recovery Limits
108	60-140
	<50 <50 <300

BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

roject#: 722450.14020 Prep Method: EPA 3520

Location: Beale AFB

BLANK SPIKE/BLANK SPIKE DUPLICATE

Matrix: Water Prep Date: 03/18/96 Batch#: 26473 Analysis Date: 03/19/96

Units: ug/L Diln Fac: 1

BS Lab ID: QC17320

Analyte	Spike Added BS	%Rec #	Limits
Diesel Range	2475 2203	89	60-140
Surrogate	%Rec Limits	3	
Hexacosane	95 60-140)	

BSD Lab ID: QC17321

Analyte	Spike Added	BSD	%Rec #	Limits	RPD #	Limit
esel Range	2475	2640	107	60-140	18	<35
burrogate	%Rec	Limi	ts		· · · · · · · · · · · · · · · · · · ·	
Hexacosane	93	60-1	40			

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

GC15 Channel A Surrogate

Sample Name : S,124796-004,26473

: C:\GC15\CHB\080B017.raw FileName

Method : DUAL

Start Time : 0.00 min Scale Factor: 0.0

End Time : 31.90 min

Plot Offset: 32 mV

Sample #: 500:2.5

Date: 3/21/96 04:23 AM

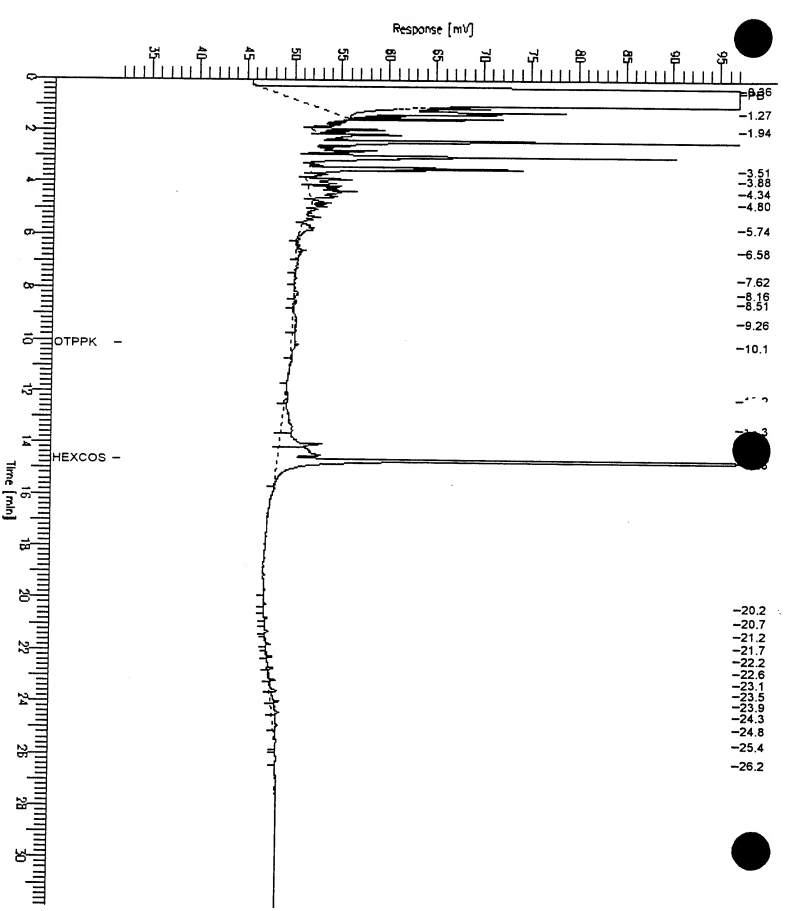
Time of Injection: 3/21/96 03:50 AM

Low Point : 32.00 mV

High Point : 97.00 mV

Page 1 of 1

Plot Scale: 65.0 mV



JP-5

GC15 Channel A Surrogate

Sample Name : CCV,96WS2068,JP5 FileName : C:\GC15\CHB\090B304.raw

Method : DUAL

Start Time : 0.00 min

End Time : 31.90 min

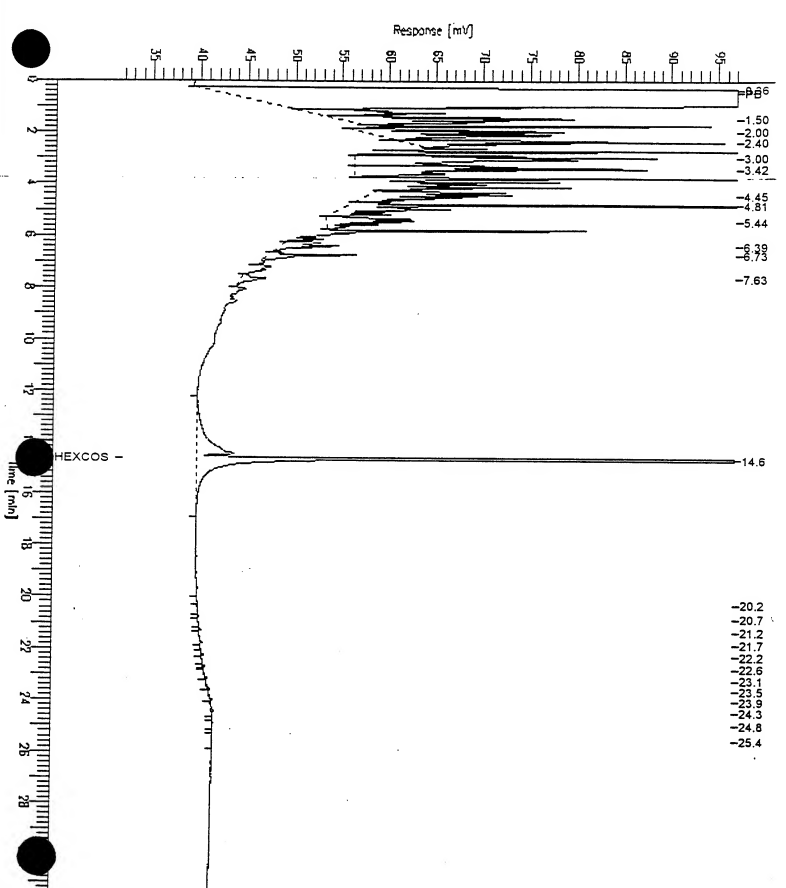
Plot Offset: 32 mV

Sample #: 250MG/L Page 1 of 1

Date: 3/20/96 06:58 PM Time of Injection: 3/20/96 06:19 PM

Low Point : 32.00 mV High Point : 97.00 mV

Plot Scale: 65.0 mV



GC15 Channel A TEH

Sample Name : CCV, 96WS1948, DSL

: C:\GC15\CHB\079B024.RAW **FileName**

: BTEH.MTH Method

Start Time : 0.01 min Scale Factor: 0.0

End Time : 19.80 min

Plot Offset: 45 mV

Sample #: 500MG/L

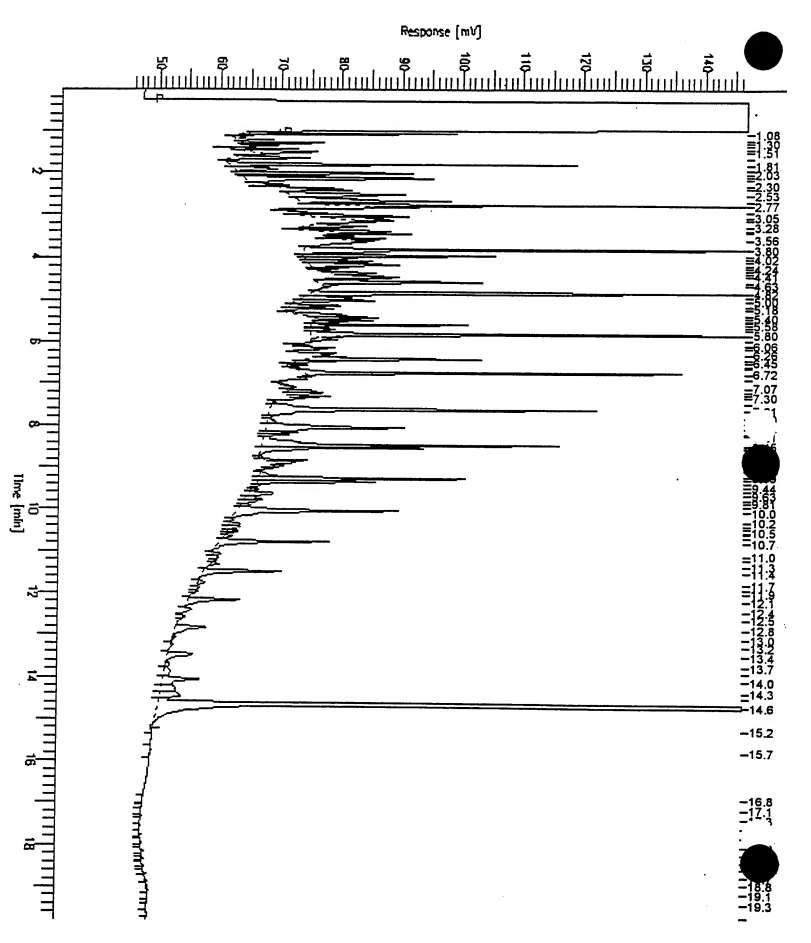
Date: 3/20/96 11:00 AM

Time of Injection: 3/20/96 12:52 AM

Low Point: 45.37 mV High Point : 146.83 mV

Page 1 of 1

Plot Scale: 101.5 mV





TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124796-001 BE-MP15-01	26573	03/11/96	03/23/96	03/23/96	
124796-002 BE-MP17-01	26573	03/11/96	03/22/96	03/22/96	
124796-003 BE-MP40-01	26573	03/11/96	03/23/96	03/23/96	
124796-004 BE-MP22-01	26573	03/12/96	03/23/96	03/23/96	

Analyte Diln Fac:	Units	124796-001 1	124796-002 1	124796-003 1	124796-004 1
Gasoline	ug/L	130 Y	<50	<50	770 Y
Surrogate			- 10 ⁻¹⁰⁴	Professional Control of Control o	
Trifluorotoluene	%REC	100	100	87	101
Promobenzene	%REC	97	97	83	106

Sample exhibits fuel pattern which does not resemble standard



TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015)

Project#: 722450.14020 Location: Beale AFB

Prep Method: EPA 5030

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124796-005	BE-MP42-01	26573	03/12/96	03/23/96	03/23/96	
124796-006	BE-MP43-01	26573	03/12/96	03/23/96	03/23/96	

Analyte Diln Fac:	Units	124796-005 1	124796-006 1	
Gasoline	ug/L	790 Y	<50	
Surrogate				
Trifluorotoluene	%REC	101	85	
Bromobenzene	%REC	105	81	

Y: Sample exhibits fuel pattern which does not resemble standard



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Water

Prep Method: EPA 5030

Location: Beale AFB

METHOD BLANK

Prep Date: 03/22/96 Analysis Date: 03/22/96

Batch#: 26573 Units: ug/L

Diln Fac: 1

Matrix:

MB Lab ID: QC17665

Analyte	Result	
Gasoline	<50	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene	98	69-120
Bromobenzene	91	70-122



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Parsons Engineering Science, Inc. Client:

Project#: 722450.14020 Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26573 Units: ug/L

Diln Fac: 1

Prep Date:

03/22/96

Analysis Date: 03/22/96

LCS Lab ID: QC17666

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	2008	2006	100	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	106	69-120		
Bromobenzene	112	70-122		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits

FileName : G:\GC05\082H033.raw

Start Time : 0.00 min Scale Factor: -1

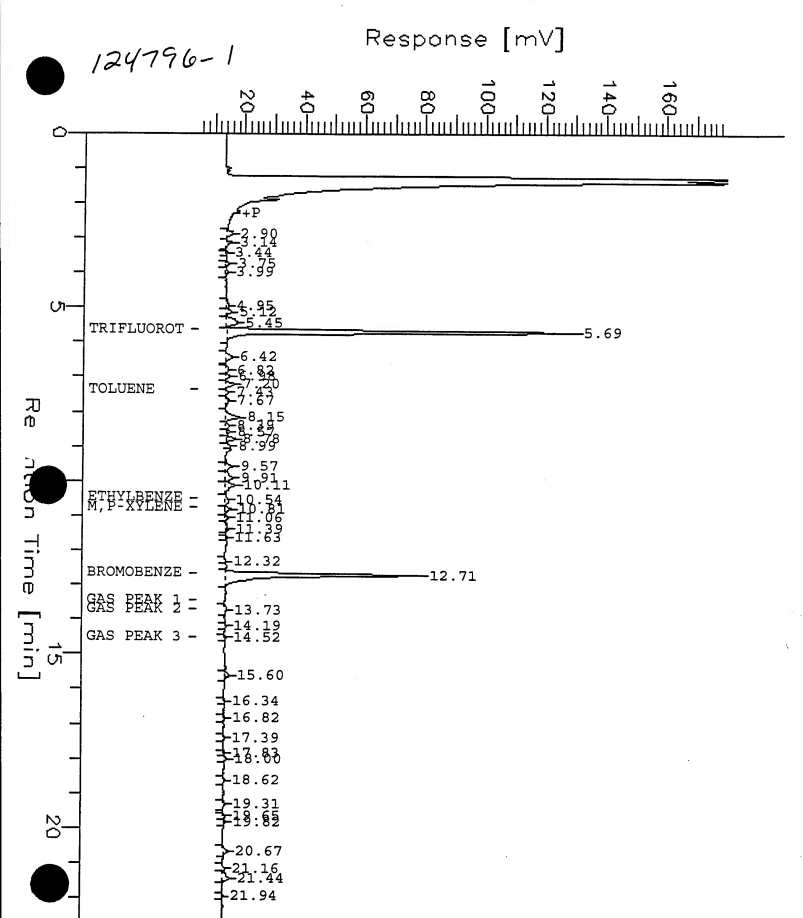
End Time : 23.42 min

Plot Offset: 5 mV

Date: 3/23/96 9:35 AM

Low Point : 4.82 mV Plot Scale: 175 mV

Page 1 of 1 High Point: 179.82 mV



FileName

: G:\GC05\082H020.raw

Start Time : 0.00 min

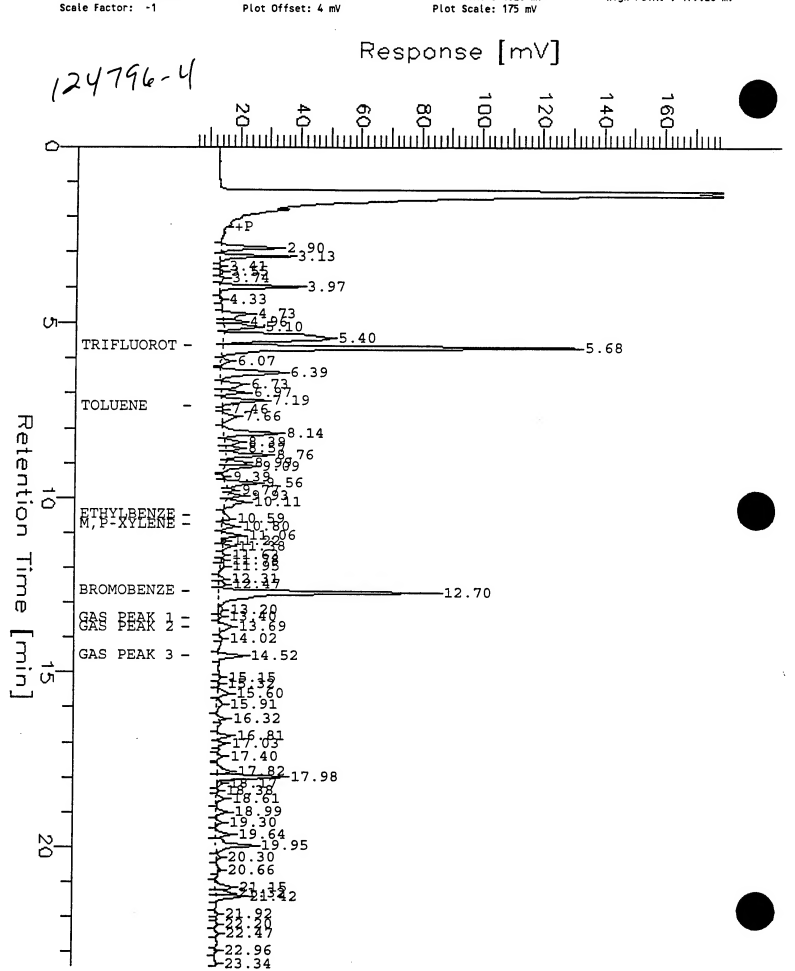
End Time : 23.42 min Plot Offset: 4 mV

Date: 3/23/96 1:29 AM

Low Point : 4.20 mV

Plot Scale: 175 mV

Page 1 of 1 High Point: 179.20 mV



FileName : G:\GC05\082H021.raw

Start Time : 0.00 min

End Time : 23.42 min

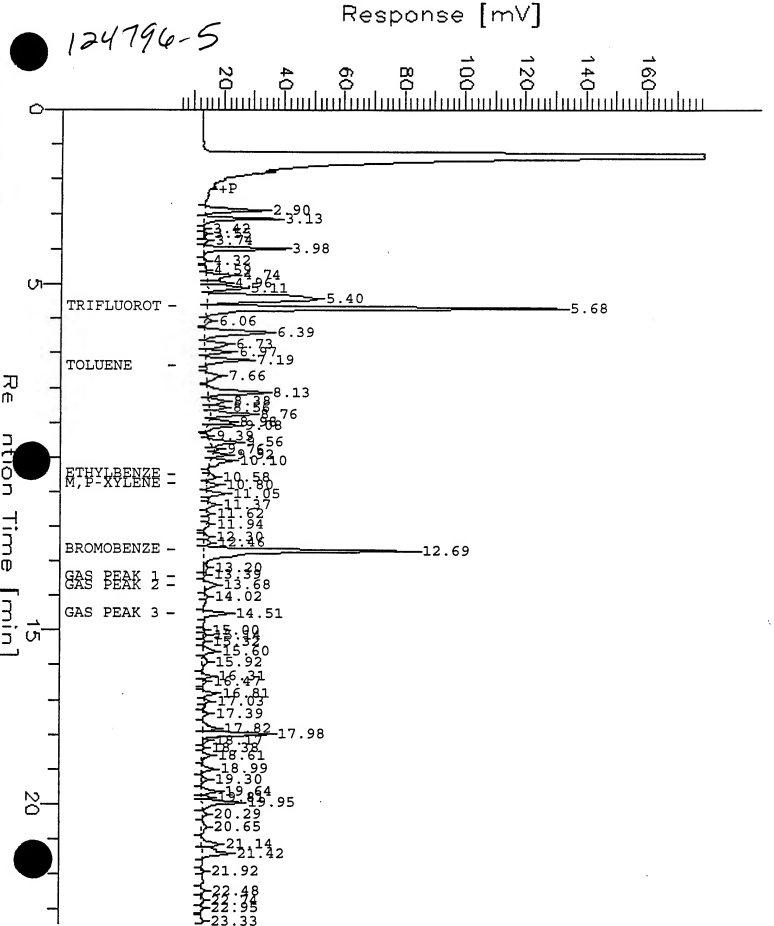
2 min Low

Date: 3/23/96 2:06 AM Low Point: 4.16 mV Page 1 of 1 High Point : 179.16 mV

Scale Factor: -1 Plot Offset: 4 mV

Plot Scale: 175 mV





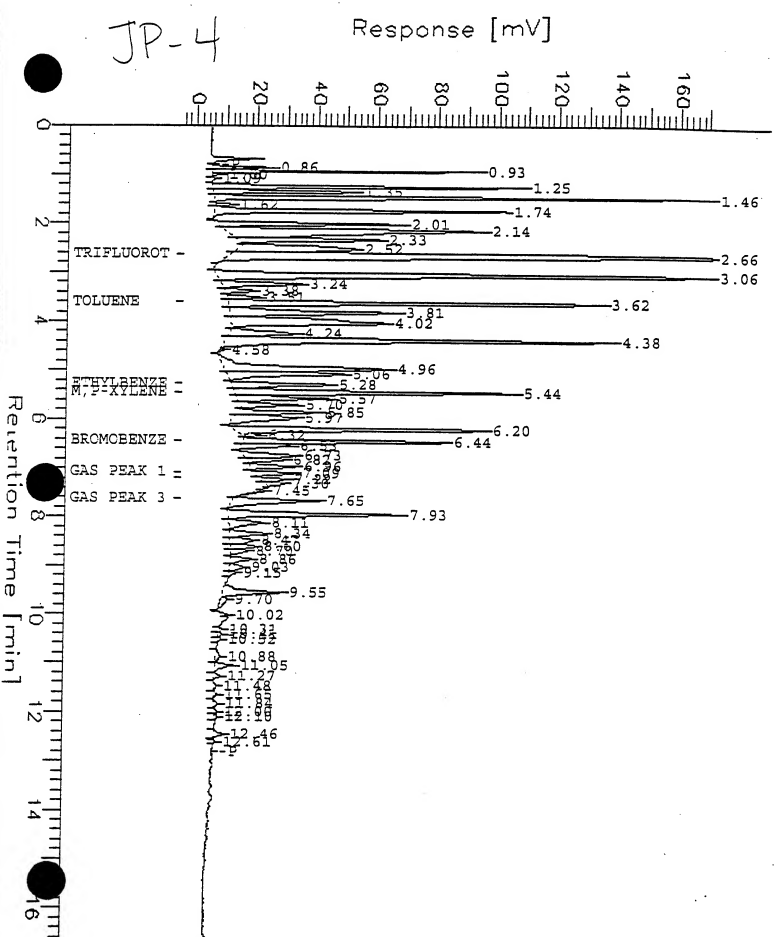
TVH2 - GC-04 RTX-1 C+T Gasoline Std. FileName : G:\GC04\135J001.raw Date : 5/15/95 9:53 AM Low Point : 42.92 mV Page 1 of 1 Start Time : 0.00 min End Time : 17.00 min High Point : 217.92 mV Scale Factor: -1 Plot Offset: 43 mV Plot Scale: 175 mV Response [mV] 200 = 1 180 = 1 160 = 1 140 = 1 -0.87 -1.15 1.35 1.60 1.83 2, 2, 03 TRIFLUOROT -2.43 2.85 -3.02 TOLUENE -3.39 3.91° -4.18 ETHYLBENZE = -5.20 -5.50 -5.96 BROMOBENZE -6.: -6.49 GAS PEAK 1 = ·6. 7.14 GAS PEAK 3 --7.40 7.70 7.87 8.06 <u>888</u>31 -9.46 10.82 11.40 11.80 11.30 12.35 1-12.77 -14.54

> 16.10 -16.45

Relention

Time

Date: 2/1/95 3:57 PM Page 1 of 1
Low Point: -4.15 mV High Point: 170.85 mV
Plot Scale: 175 mV





BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124796-001 BE-MP15-01	26573	03/11/96	03/23/96	03/23/96	
124796-002 BE-MP17-01	26573	03/11/96	03/22/96	03/22/96	
124796-003 BE-MP40-01	26573	03/11/96	03/23/96	03/23/96	
124796-004 BE-MP22-01	26573	03/12/96	03/23/96	03/23/96	

Analyte t Diln Fac:	Units	124796-001 1	124796-002 1	124796-003 1	124796-004 1
Benzene	ug/L	<0.5	<0.5	<0.5	<0.5
Toluene	ug/L	0.6	<0.5	<0.5	<0.5
Ethylbenzene	ug/L	0.7	<0.5	<0.5	<0.5
m,p-Xylenes	ug/L	0.8	<0.5	<0.5	2.2C
o-Xylene	ug/L	<0.5	<0.5	<0.5	<0.5
1,2,3-Trimethylbenzene		<0.5	<0.5	<0.5	<0.5
1,2,4-Trimethylbenzene	e ug/L	<0.5	<0.5	<0.5	<0.5
1,3,5-Trimethylbenzene	e ug/L	<0.5	<0.5	<0.5	<0.5
Surrogate					
Trifluorotoluene	%REC	99	98	86	100
Bromobenzene	%REC	90	90	76	95

C: Presence of this compound confirmed by second column, however, the confirmation concentration differed from the reported result by more than a factor of two



BTXE

lient: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124796-005 BE-MP42-01	26573	03/12/96	03/23/96	03/23/96	
124796-006 BE-MP43-01	26573	03/12/96	03/23/96	03/23/96	

Analyte U	nits	124796-005	124796-006 . 1	
Benzene	ug/L	<0.5	<0.5	
Toluene	ug/L	<0.5	<0.5	
Ethylbenzene	ug/L	1.8C	<0.5	
m,p-Xylenes	ug/L	<0.5	<0.5	
o-Xylene	ug/L	<0.5	<0.5	
1,2,3-Trimethylbenzene		<0.5	<0.5	
1,2,4-Trimethylbenzene	ug/L	<0.5	<0.5	
3,5-Trimethylbenzene	ug/L	2.2C	<0.5	
Surrogate				
Trifluorotoluene	%REC	100	84	
Bromobenzene	%REC	95	75	

C: Presence of this compound confirmed by second column, however, the confirmation concentration differed from the reported result by more than a factor of two



BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

Project#: 722450.14020

Location: Beale AFB

Prep Method: EPA 5030

METHOD BLANK

Matrix: Water Batch#: 26573 Prep Date:

03/22/96 Analysis Date: 03/22/96

Units: ug/L Diln Fac: 1

MB Lab ID: QC17665

Analyte	Result	
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	

Surrogate	%Rec	Recovery Limits
Trifluorotoluene	80	58-130
Bromobenzene	88	62-131



BATCH QC REPORT

Page 1 of 1

BTXE

client: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

Project#: 722450.14020

Prep Method:

EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Prep Date:

03/22/96

Matrix: Water Batch#: 26573 Units: ug/L

Analysis Date:

03/22/96

Diln Fac: 1

LCS Lab ID: QC17667

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	18.5	20	93	80-120
Toluene	18.4	20	92	80-120
Ethylbenzene	18	20	90	80-120
m,p-Xylenes	36.6	40	92	80-120
o-Xylene	18.5	20	93	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	100	58-130		
Bromobenzene	90	62-131		

Column to be used to flag recovery and RPD values with an asterisk alues outside of QC limits

ke Recovery: O out of 5 outside limits

EPA 5030

Lab #: 124796

BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc. Analysis Method: EPA 8020 Project#: 722450.14020 Prep Method:

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Sample Date: 03/19/96 Lab ID: 124900-009 Received Date: 03/20/96 Matrix: Water Prep Date: 03/22/96 Batch#: 26573 Analysis Date: 03/22/96

Units: ug/L Diln Fac: 1

MS Lab ID: QC17668

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Benzene	20	<0.5000	19.1	96	75-125
Toluene	20	<0.5000	19.1	96	75-125
Ethylbenzene	20	<0.5000	19.1	96	75-125
m,p-Xylenes	40	<0.5000	38.1	95	75-125
o-Xylene	20	<0.5000	19.4	97	75-125
Surrogate	%Rec	Limits			
Trifluorotoluene	101	58-130			
Bromobenzene	91	62-131			

MSD Lab ID: QC17669

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Benzene	20	19.8	99	75-125	4	<20
Toluene	20	19.7	99	75-125	3	<20
Ethylbenzene	20	19.5	98	75-125	2	<20
m,p-Xylenes	40	39	98	75-125	2	<20
o-Xylene	20	19.8	99	75-125	2	<20
Surrogate	%Rec	Limits	3			
Trifluorotoluene	101	58-130)	101		
Bromobenzene	94	62-13:				

Column to be used to flag recovery and RPD values with an asterisk

* Values outside of QC limits RPD: 0 out of 5 outside limits

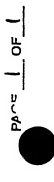
Spike Recovery: 0 out of 10 outside limits



1301 Marina Village Pkwy, Suite 200 Alameda, California 94501 Phone: (510) 769-0100 FAX: (510) 769-9244







Project No.: 722450.14020								٥	OFIGANICS	SS			INORG	NORGANICS		/отнея/
Project Name/Location: Beale AFB, UST Site	UST SIE	6 1					-	١٥/	/	/	/		//	_		
Project Manager: Todd Wiedemeier (Denver, CO)	(Denver,	8				- 4	16	39700	100	90	\	<u> </u>		<u> </u>		<u>'</u>
Site Manager: Michael Phelps (Alameda, CA)	eda, CA)					NE	200	100 (U	00 00	/	\	_	<u> </u>	6		<u> </u>
Sampler(s): (Initials and Signature(s))	-					W S	31	105 105 105	0108	\	\	\	\	122-Y		
MO P. Janes A Grant	win	\				THE STATE OF THE S	08/0	08/06	υεος		(8)	E5.3)		(SO)		_
TATE - Melanger	- }- }	1			O : OF (JANA SOEOZI OF SOEOZI HAT	HOL	Was (SQ)	(0Z58/0S	(e	A AIGH W	ESI OIO		DWIST		Note: All analyses and holding times
Sample ID Date	Time	Begin Depth	End	Matrix		TOSEB-HO	PURE PLANT	W (E32) NO (9080) OC (9080	2000	W (E35).	Iselinity (3	9) 4011	A2/0	Sein-eize		according to AFCEE handbook requirements.
BE-MP15-41 3/11/96	0401	1	1	soil	ב	3 7	8 7	6 3		V	थ	0	4.	干		Am a con 1
2 BE-MP17-01 3/11/96	1230	ı	(Soil	メ	×	×				+-				5	
3 BE-MP44-01 3/4/96	१०५१	Ì	1	ligs	× ×	J	7									
4 BE- MP22-01 3/12/96 1120	0211	1	(llos	X ک	7.	×							-		
5 BE-MP42-B1 3/12/96 1130	1130		1	lios /	7	×	У									
16 BE-MP43-61 3/12/96	0711	١	/	soil	3 4											-3
				soil												
	ממן,	ala	2,77	4 soil									1			
			1/2	soil												
				soll							-				7	DV CCV
				soil					17		-					9 20
				soil												15/16
				soil							-					
11/2			EIVED BY:	HEGEIVED BY: (SIGNATUR [WALLL: CORP)	18E) / (3)	2 - 9 E	NOTES: 1. Secondary column confirmation required. 2. Quantify all carbon ranges: diesel, jet fuel, oll/grease. 3. Delonized water extraction	y column ill carbon water ex	confirm ranges: traction	ation red diesel, j	ulred. et fuel,	Morena	1.	4. TMB	r. 1,2,3-t trimet	4. TMBs: 1,2,3-trimethylbenzene, 1,2,4, trimethylbenzene, and 1,3,5-trimethylbenzene
neundoished BY: (SignAlOHE) DAIE		IIME PECE	EIVED FOR	LAB BY:((§IGNATÙRE)		7E	TIME	REMAR	REMARKS/COMMENTS:	MENTS					
						-										

Rev. 1 4/95

	4: 124796 Date Received: 3/13/46 Number of Coolers:
A.	Preliminary Examination Phase
	Date Opened: 3/13/4 6 By (print): 4 thia cho(sign) (Stue C Solle)
1.	Did cooler come with a shipping slip (airbill, etc.)?
•	If YES, enter carrier name and airbill number: FED BX Were custody seals on outside of cooler? YES NO
2.	// 010 000000 000 000 000 000 00 000 00 00
2	How many and where? Seal date: Seal name: Were custody seals unbroken and intact at the date and time of arrival? YES NO
3.	Were custody seals unbroken and intact at the date and time of arrival?
4.	Were custody papers dry and intact when received?
5.	Were custody papers filled out properly (ink, signed, etc.)?
6.	Did you sign the custody papers in the appropriate place?
7.	Was project identifiable from custody papers?
•	If YES, enter project name at the top of this form. If required, was sufficient ice used? YES NO
8.	11.10(4 02, 11
	Type of ice: 1ext Temperature: 40C
ח	Lagin Phase
B.	Date Logged In: 3/13/96 By (print): Cynthia Sella (sign) Cynthia Sella
1	Describe time of posking in cooler: See to had a see to had
1. 2.	Describe type of packing in cooler: paper \$ bubble lings Did all bottles arrive unbroken? (ES) NO
2. 3.	Were labels in good condition and complete (ID, date, time, signature, etc.)? (YES) NO
<i>3</i> .	Did bottle labels agree with custody papers?
4 . 5.	Were appropriate containers used for the tests indicated? YES' NO
5. 6.	Were correct preservatives added to samples?
7 .	Was sufficient amount of sample sent for tests indicated?
7 . 3 .	Were bubbles absent in VOA samples? If NO, list sample Ids belowYES NO
9.	Was the client contacted concerning this sample delivery? YES NO
7 .	If YES, give details below.
	Who was called? M. Phelos By whom? The Date: 3/13/96
/ ,	Willo was caned. 19(5) Telloss 2) Williams 1950 2000.
Additio	onal Comments:
, ladini	, E
	DE-meis-01 did have a comber 16
	michael Philos 1- formed 3/13/96. Them
	The same of the sa
	<u> </u>

Filename: F:\qc\forms\cooler.wpd



Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 94710, Phone (510) 486-0900

ANALYTICAL REPORT

Prepared for:

Parsons Engineering Science, Inc. 1301 Marina Village Parkway Suite 200 Alameda, CA 94501

Date: 02-APR-96

Lab Job Number: 124797

Project ID: 722450.14020 Location: Beale AFB

Reviewed by:

Reviewed by:

This package may be reproduced only in its entirety.

moe K Morris

MEMORANDUM TO FILE

•	JOB NO. 722450. 14020
·*	FILE DESIGNATION Beale AFB-UST Site
	DATE 4/5/96 TIME
PHONE CALL FROM Michael Phelps	
PHONE CALL TO	PHONE NO
CONFERENCE WITH	
	<i>(</i>
SUBJECT Incorrect Sample Lesignation	on on COC dated 3/12/96
The sample disignated BE-1	
Lesignated BE-MAPP-01. There is	vas no sample collected from
MP-\$8.	
•	
	Menl Alle



Laboratory Number: 124797

Client: Parsons Engineering Science

Project#: 722450.14020 Location: Beale AFB Sample Date: 03/12/96 Receipt Date: 03/13/96

CASE NARRATIVE

This hardcopy data package contains sample results and batch QC for six water samples received from the above referenced project on March 13, 1996. All samples were received cold and intact.

TPH/Extractable: No analytical problems were encountered.

TPH/Purgeable: Due to analyst oversight, sample BE-MP46-01 (C&T# 124797-006) was not analyzed until one day past the technical holding time. Sample chromatograms do not resemble gasoline, but are similar to JP-4.

BTXE (EPA 8020): Due to analyst oversight, sample BE-MP46-01 (C&T# 124797-006) was not analyzed until one day past the technical holding time.



TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 3520

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124797-001 BE-MP11-01	26513	03/12/96	03/19/96	03/22/96	
124797-002 BE-MP44-01	26513	03/12/96	03/19/96	03/22/96	
124797-004 BE-MP08-01	26513	03/12/96	03/19/96	03/22/96	
124797-005 BE-MP41-01	26513	03/12/96	03/19/96	03/22/96	

Analyte Diln Fac:	Units	124797-001 1		124797-002 1		124797-004 1		124797-005 1	
JP5	ug/L	130	Y	190	Y	710	YH	700	YH
Diesel Range	ug/L	<50		94	\mathtt{YL}	4500	YH	4500	YH
Motor Oil Range	ug/L	<300		<300		3500	Y	3800	Y
Surrogate		100						*	
Hexacosane	*REC	111		106		78		81	

Y: Sample exhibits fuel pattern which does not resemble standard

H: Heavier hydrocarbons than indicated standard

L: Lighter hydrocarbons than indicated standard



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 3520

Location: Beale AFB

METHOD BLANK

Prep Date: 03/19/96

Matrix: Water Batch#: 26513 Units: ug/L Diln Fac: 1

Analysis Date: 03/21/96

MB Lab ID: QC17466

Analyte	Result	
JP5	<50	
Diesel Range	<50	
Motor Oil Range	<300	
Surrogate	%Rec	Recovery Limits
Hexacosane	103	60-140



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method:

EPA 3520

BLANK SPIKE/BLANK SPIKE DUPLICATE

Matrix: Water Batch#: 26513 Units: ug/L Diln Fac: 1

Prep Date: Analysis Date: 03/19/96 03/21/96

BS Lab ID: QC17467

Analyte	Spike Added	BS	%Rec #	Limits
Diesel Range	2475 2	218	90	60-140
Surrogate	%Rec	Limits		
Hexacosane	96	60-140		

BSD Lab ID: QC17468

Analyte	Spike Added	BSD	%Rec #	Limits	RPD #	Limit
Diesel Range	2475	2330	94	60-140	5	<35
Surrogate	%Rec	Limit	s			
Hexacosane	95	60-14	0			

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

GC15 Channel A Surrogate

Sample Name : S, 124797-001, 26513 FileName : C:\GC15\CHB\080B064.raw

: DUAL Method

Start Time : 0.00 min Scale Factor: 0.0

End Time : 31.90 min Plot Offset: 32 mV

Sample #: 500:2.5

Date: 3/22/96 02:54 PM

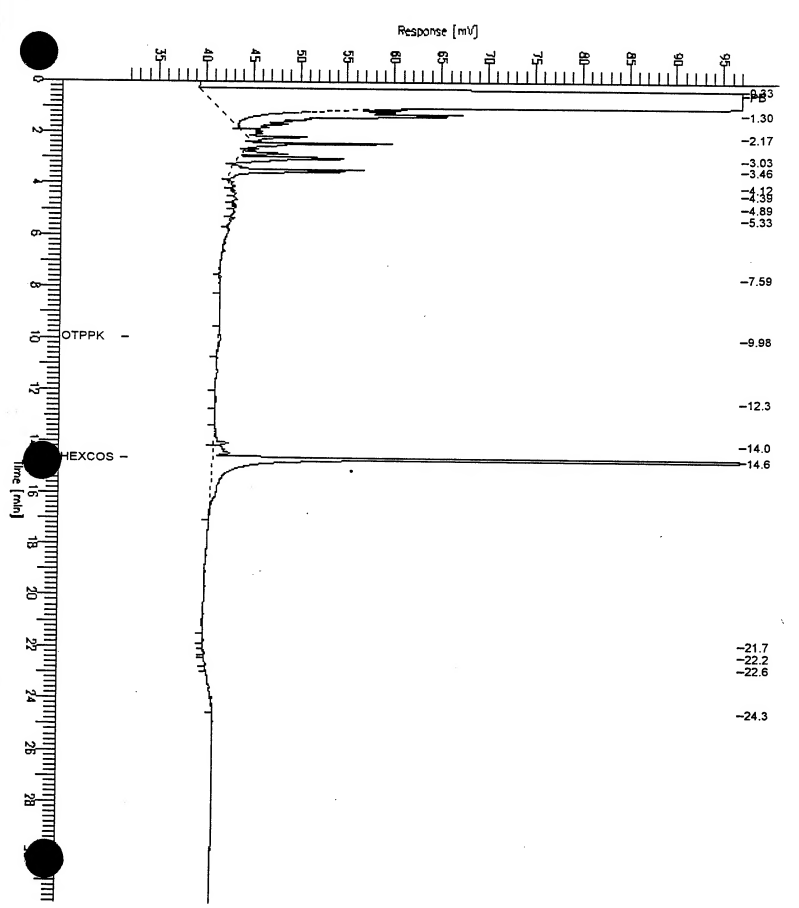
Time of Injection: 3/22/96 02:21 PM

Low Point : 32.00 mV

High Point : 97.00 mV

Page 1 of 1

Plot Scale: 65.0 mV



CHAIMEL Durroyace

Sample Name : S,124797-002,26513 FileName : C:\GC15\CHB\080B059.raw

Method : DUAL

Scale Factor: 0.0

Start Time : 0.00 min

End Time : 31.90 min Plot Offset: 32 mV

Sample #: 500:2.5

Date: 3/22/96 11:07 AM

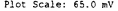
Time of Injection: 3/22/96 10:34 AM

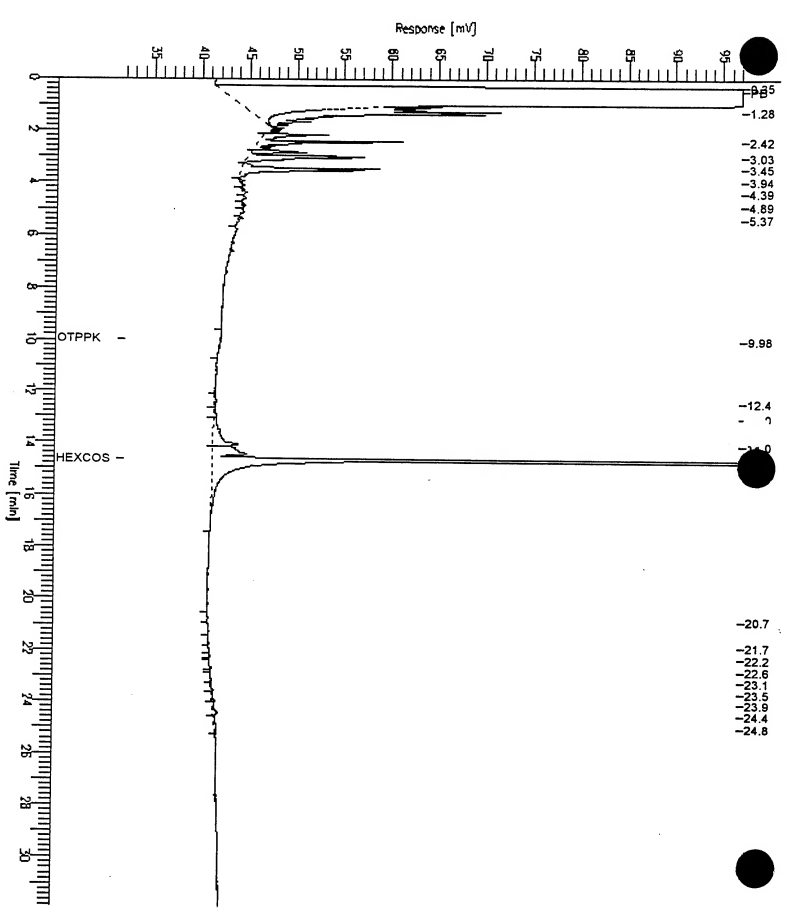
Low Point : 32.00 mV

Plot Scale: 65.0 mV

High Point : 97.00 mV

Page 1 of 1





GC15 Channel A TEH

Sample Name : S,124797-004,26513 : C:\GC15\CHB\080B060.RAW FileName

Method : BTEH.MTH

Start Time : 0.01 min Sr 'a Factor: 0.0

End Time : 31.91 min

Plot Offset: 38 mV

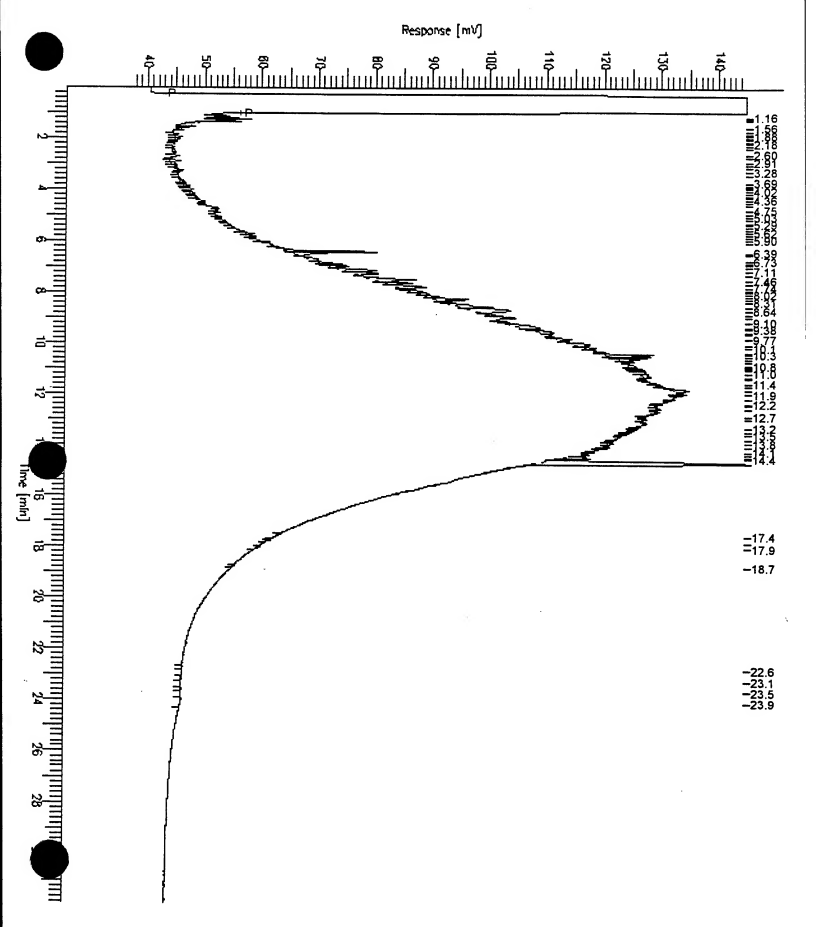
Sample #: 500:2.5 Date: 3/25/96 04:47 PM

Time of Injection: 3/22/96 11:21 AM

Low Point : 37.75 mV High Point : 144.96 mV

Page 1 of 1

Plot Scale: 107.2 mV



Sample Name: S, 124797-005, 26513 FileName : C:\GC15\CHB\080B062.RAW

: BSURR.MTH Method

Start Time : 0.01 min Scale Factor: 0.0

End Time : 31.91 min Plot Offset: 39 mV

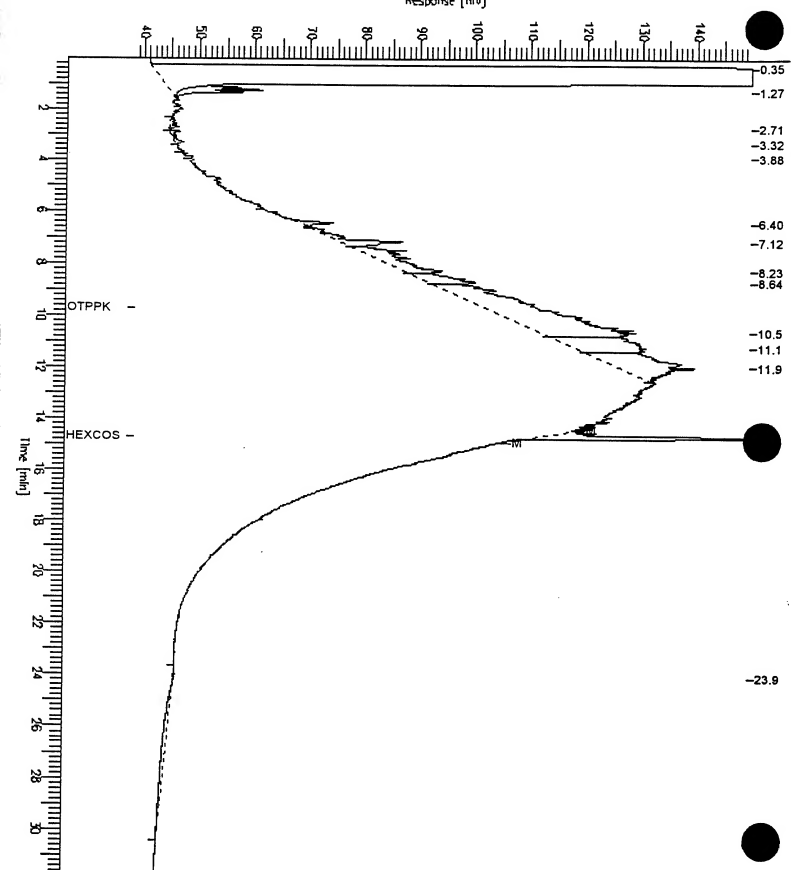
Sample #: 500:2.5 Page 1 of 1 Date: 3/25/96 04:52 PM

Time of Injection: 3/22/96 12:53 PM

Low Point : 38.58 mV High Point: 149.83 mV

Plot Scale: 111.3 mV





JR.S

GC15 Channel A TEH

Sample Name : CCV, 96WS2068, JP5

FileName : C:\GC15\CHB\080B050.RAW

Method : BTEH.MTH Start Time : 0.01 min

Factor: 0.0

End Time : 31.91 min Plot Offset: 43 mV

Sample #: 250MG/L

Date: 3/25/96 05:13 PM

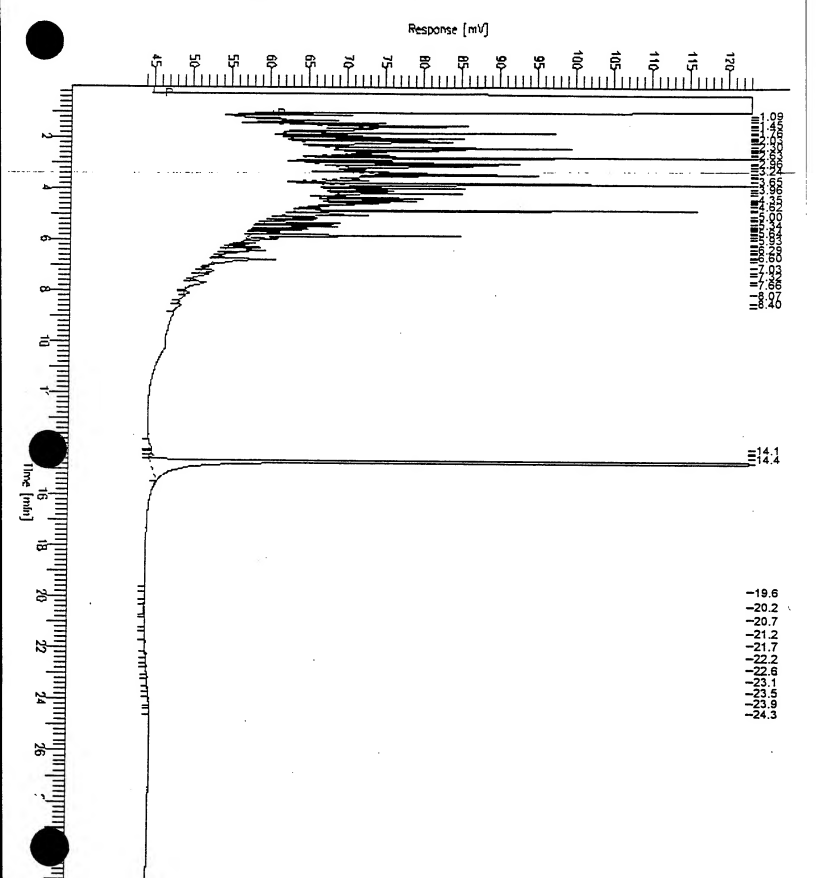
Time of Injection: 3/22/96 04:00 AM

Low Point : 43.16 mV

High Point : 123.07 mV

Page 1 of 1

Plot Scale: 79.9 mV



DIESEL

GC15 Channel A TEH

Sample Name : CCV,96WS1948,DSL FileName : C:\GC15\CHB\080B047.RAW

Method : BTEH.MTH

Start Time : 0.31 min

Scale Factor: 0.0 End Time : 31.91 min

Plot Offset: 44 mV

Sample #: 500MG/L

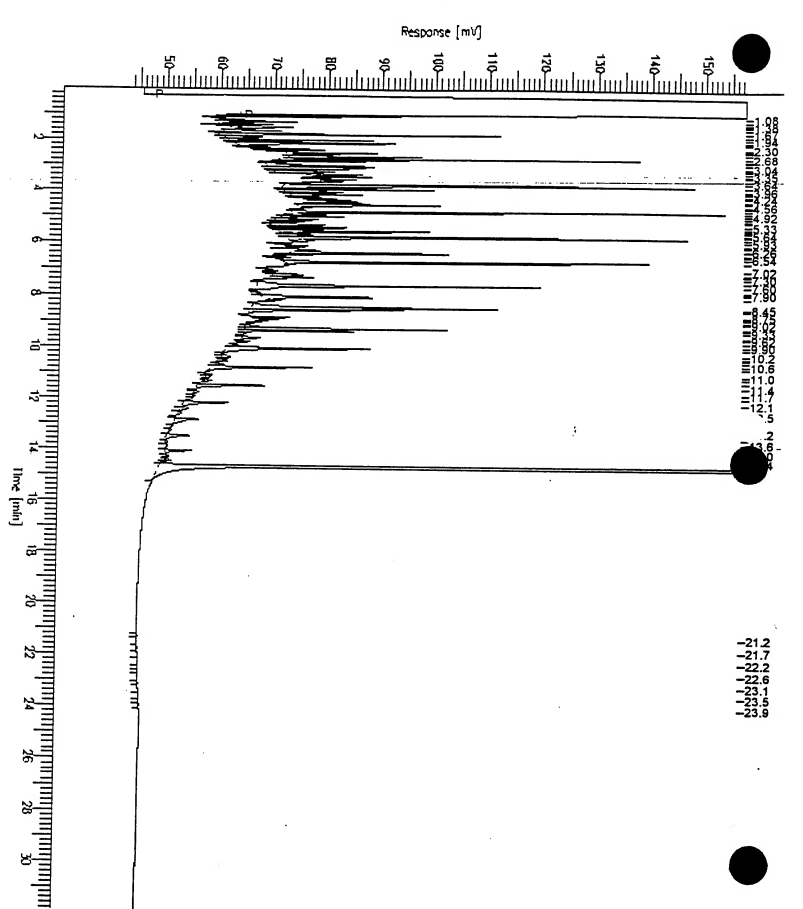
Date: 3/25/96 05:16 PM

Time of Injection: 3/22/96 01:48 AM

Low Point : 43.61 mV High Point : 157.43 mV

Page 1 of 1

Plot Scale: 113.8 mV



GC15 Channel A TEH motor Oil

Sample Name : CCV, 95WS1596, MO

FileName : C:\GC15\CHB\080B051.RAW

0.0

Methca : BTEH.MTH Start Time : 0.01 min

Pactor:

End Time : 31.91 min

Plot Offset: 41 mV

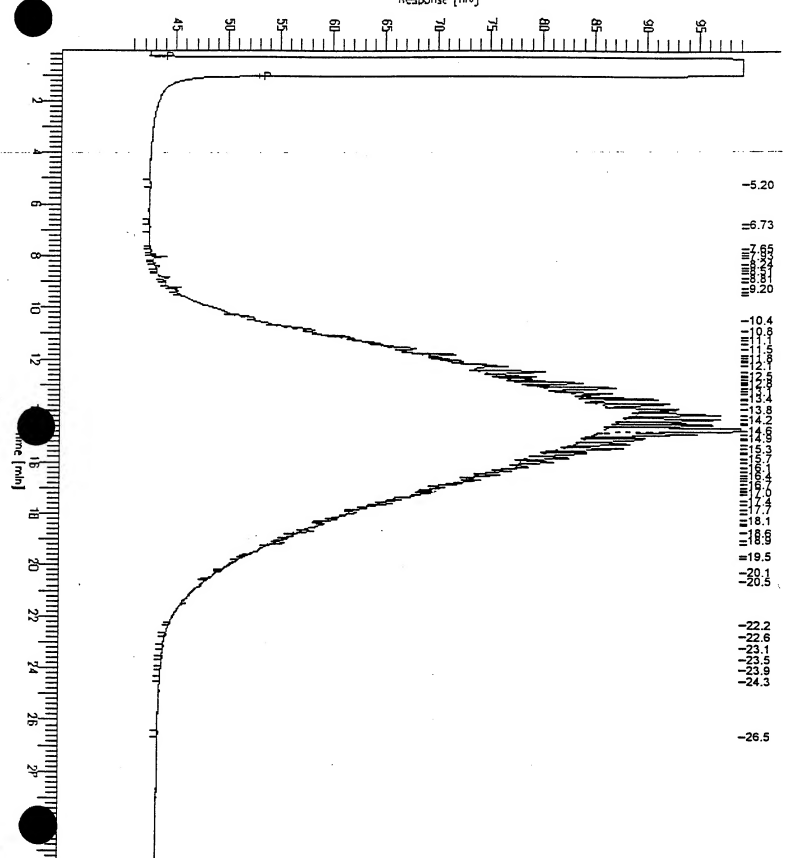
Sample #: 500MG/L

Low Point : 40.69 mV High Point : 99.19 mV

Page 1 of 1

Plot Scale: 58.5 mV







TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

•

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method:

EPA 5030

Location: Beale AFB

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124797-001 BE-MP11-01	26573	03/12/96	03/22/96	03/22/96	
124797-002 BE-MP44-01	26573	03/12/96	03/23/96	03/23/96	
124797-003 BE-MP45-01	26573	03/12/96	03/23/96	03/23/96	
124797-004 BE-MP08-01	26573	03/12/96	03/23/96	03/23/96	

Analyte Diln Fac:	Units	124797-001 1	124797-002 1	124797-003 1	124797-00 1
Gasoline	ug/L	380 Y	410 Y	<50	<50
Surrogate					
Trifluorotoluene	%REC	99	100	99	100
Bromobenzene	%REC	96	98	95	97

Y: Sample exhibits fuel pattern which does not resemble standard



TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

roject#: 722450.14020 Location: Beale AFB

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124797-005 BE-MP41-01	26573	03/12/96	03/22/96	03/22/96	
124797-006 BE-MP46-01	26672	03/12/96	03/27/96	03/27/96	

Analyte Diln Fac:	Units	124797-005 1	124797-006 1	
Gasoline	ug/L	<50	<50	
Surrogate				
Trifluorotoluene	%REC	100	96	
Bromobenzene	%REC	95	94	



BATCH QC REPORT

Page 1 of '

		TVH-Tota	l Volatil	Le Hydrocarbons	
Project#:	Parsons Engineering 722450.14020 Beale AFB	Science,	Inc.	Analysis Method: Prep Method:	CA LUFT (EPA 8015) EPA 5030
			METHOD	BLANK	
Matrix: Batch#: Units: Diln Fac:	Water 26573 ug/L 1			Prep Date: Analysis Date:	03/22/96 03/22/96

MB Lab ID: QC17665

Analyte	Result	
Gasoline	<50	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene	98	69-120
Bromobenzene	91	70-122



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

METHOD BLANK

Matrix: Water Batch#: 26672

Prep Date: 03/27/96 Analysis Date: 03/27/96

Units: ug/L Diln Fac: 1

MB Lab ID: QC18055

Analyte	Result	
Gasoline	<50	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene Bromobenzene	95 87	69-120 70-122



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26573 Units: ug/L Diln Fac: 1

Prep Date: 03/22/96 Analysis Date: 03/22/96

LCS Lab ID: QC17666

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	2008	2006	100	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	106 112	69-120 70-122		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

client: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Method: EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

 Matrix:
 Water
 Prep Date:
 03/27/96

 Batch#:
 26672
 Analysis Date:
 03/27/96

Batch#: 26672 Analys
Units: ug/L
Diln Fac: 1

LCS Lab ID: QC18056

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	1843	2006	92	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	103 101	69-120 70-122		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

pike Recovery: 0 out of 1 outside limits

BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: EPA 5030

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: BE-MP46-01 Lab ID: 124797-006 Matrix: Water Batch#: 26672

Sample Date: 03/12/96 Received Date: 03/13/96 Prep Date: Analysis Date:

03/27/96 03/27/96

Units: ug/L Diln Fac: 1

MS Lab ID: QC18058

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Gasoline	2000	<50.00	2047	102	75-125
Surrogate	%Rec	Limits			
Trifluorotoluene Bromobenzene	104 110	69-120 70-122			

MSD Lab ID: QC18059

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Gasoline	2000	2135	107	75-125	4	<20
Surrogate	%Rec	Limit	s			
Trifluorotoluene Bromobenzene	104 114	69-120 70-12	-			**

[#] Column to be used to flag recovery and RPD values with an asterisk

RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits

FileName : G:\GC05\082H010.raw

Start Time : 0.00 min

Scale Factor: -1

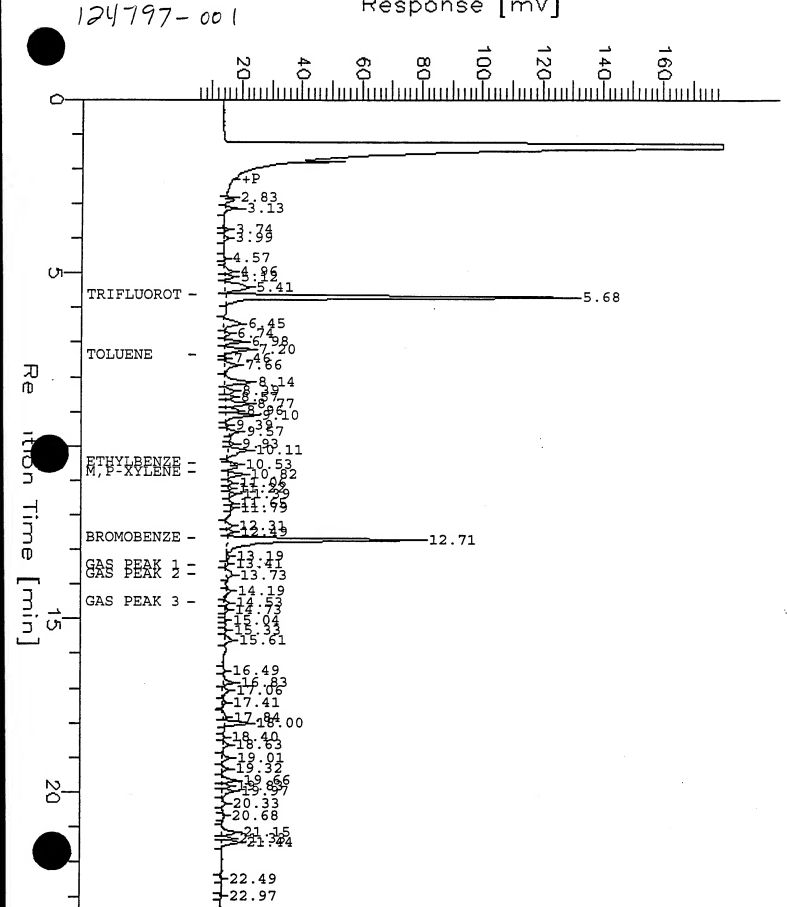
End Time : 23.42 min Plot Offset: 5 mV

Date: 3/22/96 7:13 PM Low Point : 4.96 mV

Plot Scale: 175 mV

Page 1 of 1 High Point: 179.96 mV

Response [mV]



FileName

Scale Factor: -1

: G:\GC05\082H023.raw Start Time : 0.00 min

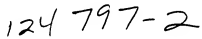
End Time : 23.42 min Plot Offset: 4 mV

Date: 3/23/96 3:21 AM

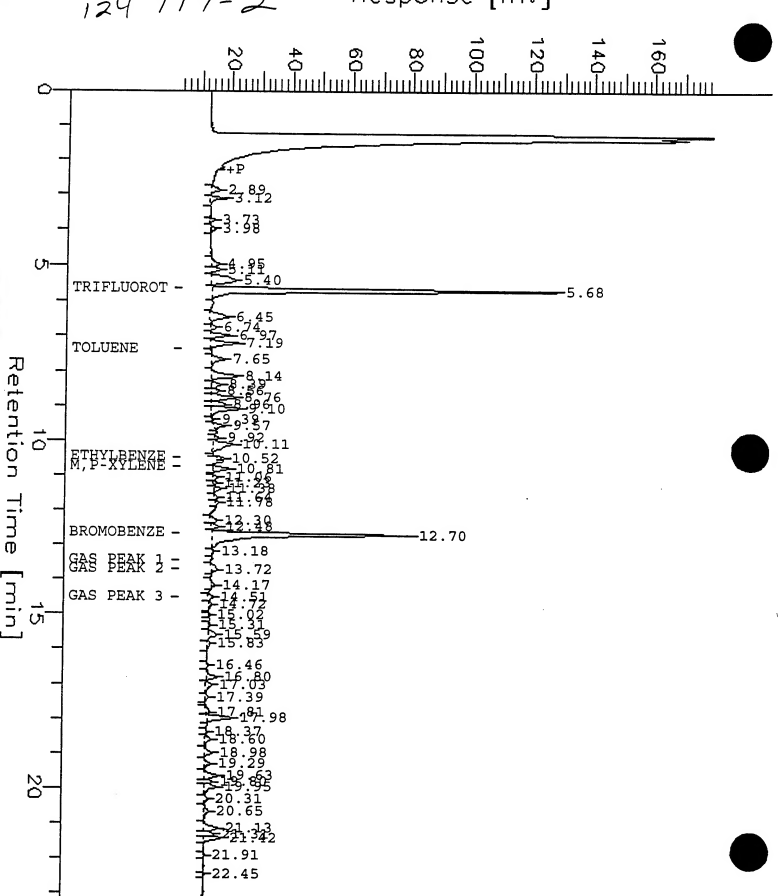
Low Point : 3.95 mV

Plot Scale: 175 mV

Page 1 of 1 High Point: 178.95 mV



Response [mV]



TVH2 - GC-04 RTX-1

C+T Gasoline Std.

: G:\GC04\135J001.raw FileName

Start Time : 0.00 min Scale Factor: -1

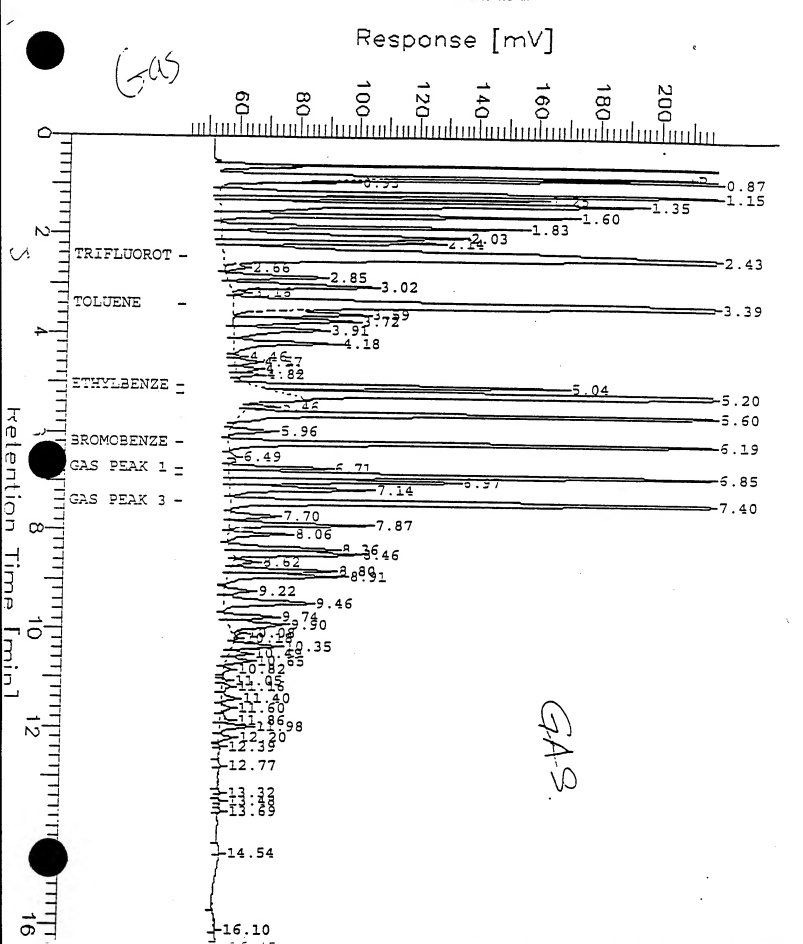
End Time : 17.00 min

Plot Offset: 43 mV

Date: 5/15/95 9:53 AM Low Point : 42.92 mV

Plot Scale: 175 mV

Page 1 of 1 High Point : 217.92 mV



1VH2 - GC-04 RTX-1 : G:\GC04\B\026J011.raw Date: 2/1/95 3:57 PM Start Time : 0.00 min Page 1 of 1 End Time : 17.00 min Low Point : -4.15 mV High Point : 170.85 mV Scale Factor: -1 Plot Offset: -4 mV Plot Scale: 175 mV Response [mV] 0.93 -1.25 1.46 -1.74 -2.14 TRIFLUOROT --2.66 -3.06 TOLUENE -3.62 $\frac{-3.81}{4.02}$ -4.38 4.58 MYPYXPENNE = -5.44 . 70⁰ . 5 / . 97 85 BROMOBENZE -GAS PEAK 1 = GAS PEAK 3 --7.65 -7.93 10.02

Retention



BTXE

client: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124797-001 BE-MP11-01	26573	03/12/96	03/22/96	03/22/96	
124797-002 BE-MP44-01	26573	03/12/96	03/23/96	03/23/96	
124797-003 BE-MP45-01	26573	03/12/96	03/23/96	03/23/96	
124797-004 BE-MP08-01	26573	03/12/96	03/23/96	03/23/96	

Analyte t Diln Fac:	Inits	124797-001 1	124797-002 1	124797-003 1	124797 - 004 1
Benzene	ug/L	<0.5	<0.5	<0.5	<0.5
Toluene	ug/L	<0.5	<0.5	<0.5	<0.5
Ethylbenzene	ug/L	<0.5	<0.5	<0.5	<0.5
m,p-Xylenes	ug/L	<0.5	<0.5	<0.5	<0.5
o-Xylene	ug/L	<0.5	<0.5	<0.5	<0.5
,2,3-Trimethylbenzene	ug/L	<0.5	1.4	<0.5	<0.5
,2,4-Trimethylbenzene	ug/L	<0.5	<0.5	<0.5	<0.5
1,3,5-Trimethylbenzene	ug/L	<0.5	<0.5	<0.5	<0.5
Surrogate	· · · · · · · · · · · · · · · · · · ·				
Trifluorotoluene	%REC	98	99	98	80
Bromobenzene	%REC	91	91	88	84



BTXE

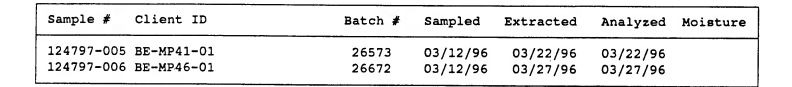
Parsons Engineering Science, Inc. Client:

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030



Analyte Diln Fac:	Units	124797-005 1	124797-006 1	
Benzene	ug/L	<0.5	<0.5	
Toluene	ug/L	<0.5	<0.5	
Ethylbenzene	ug/L	<0.5	<0.5	
m,p-Xylenes	ug/L	<0.5	<0.5	
o-Xylene	ug/L	<0.5	<0.5	
1,2,3-Trimethylbenzene	e ug/L	<0.5	<0.5	
1,2,4-Trimethylbenzene	∍ ug/L	<0.5	<0.5	
1,3,5-Trimethylbenzene	e ug/L	<0.5	<0.5	
Surrogate				
Trifluorotoluene	%REC	98	97	
Bromobenzene	%REC	89	88	



BATCH QC REPORT

Page 1 of 1

BTXE

client: Parsons Engineering Science, Inc.

30 July 3 V. S. C. V.

Analysis Method: EPA 8020

Project#: 722450.14020 Location: Beale AFB

Prep Method:

EPA 5030

METHOD BLANK

03/22/96

Matrix: Water Batch#: 26573

Prep Date: Analysis Date: 03/22/96

Units: ug/L Diln Fac: 1

MB Lab ID: QC17665

Analyte	Result	
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	

Surrogate	%Rec	Recovery Limits
rifluorotoluene	80	58-130
romobenzene	88	62-131



BATCH QC REPORT

Page 1 of 1

			1490 1 01 .
		BTXE	
Project#:	Parsons Engineering Science, In 722450.14020 Beale AFB	-	EPA 8020 EPA 5030
	M	ETHOD BLANK	
Matrix: Batch#: Units: Diln Fac:	Water 26672 ug/L 1	Prep Date: Analysis Date:	03/27/96 03/27/96

MB Lab ID: QC18055

Analyte	Result	
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene	94	58-130
Bromobenzene	81	62-131

BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26573 Units: ug/L

Diln Fac: 1

Prep Date: 03/22/96

Analysis Date: 03/22/96

LCS Lab ID: QC17667

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	18.5	20	93	80-120
Toluene	18.4	20	92	80-120
Ethylbenzene	18	20	90	80-120
m,p-Xylenes	36.6	40	92	80-120
o-Xylene	18.5	20	93	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	100	58-130		
Bromobenzene	90	62-131		

Column to be used to flag recovery and RPD values with an asterisk alues outside of QC limits

ke Recovery: 0 out of 5 outside limits



BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26672 Units: ug/L Diln Fac: 1

Prep Date: 03/27/96 Analysis Date:

03/27/96

LCS Lab ID: QC18057

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	16.8	20	84	80-120
Toluene	17.7	20	89	80-120
Ethylbenzene	16.2	20	81	80-120
m,p-Xylenes	36.8	40	92	80-120
o-Xylene	17.5	20	88	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	96	58-130		
Bromobenzene	81	62-131		
Bromobenzene				

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 5 outside limits



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plient: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ

Lab ID: 124900-009 Matrix: Water

Batch#: 26573 Units: ug/L Diln Fac: 1

Sample Date:

03/19/96 Received Date: 03/20/96 Prep Date: 03/22/96

Analysis Date: 03/22/96

MS Lab ID: QC17668

Analyte	Spike Added	Sample	MS	%Rec #	Limits	
Benzene	20	<0.5000	19.1	96	75-125	
Toluene	20	<0.5000	19.1	96	75-125	
Ethylbenzene	20	<0.5000	19.1	96	75-125	
m,p-Xylenes	40	<0.5000	38.1	95	75-125	
o-Xylene	20	<0.5000	19.4	97	75-125	
Surrogate	%Rec	Limits				
Trifluorotoluene	101	58-130				
Bromobenzene .	91	62-131				

MSD Lab ID: QC17669

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Benzene	20	19.8	99	75-125	4	<20
Toluene	20	19.7	99	75-125	3	<20
Ethylbenzene	20	19.5	98	75-125	2	<20
m,p-Xylenes	40	39	98	75-125	2	<20
o-Xylene	20	19.8	99	75-125	2	<20
Surrogate	%Rec	Limit	s			
Trifluorotoluene	101	58-13	0		 	
Bromobenzene	94	62-13	1			

[#] Column to be used to flag recovery and RPD values with an asterisk

Spike Recovery: 0 out of 10 outside limits

^{*} Values outside of QC limits RPD: 0 out of 5 outside limits

PARSONS ENGINEERING SCIENCE, INC.

1301 Marina Village Pkwy, Suite 200 Alameda, California 94501

Phone: (510) 769-0100 FAX: (510) 769-9244

CHAIN OF CUSTODY RECORD

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OF \

PAGE 1

handbook requirements. according to AFCEE and holding times Note: All analyses Remarks OTHER INOFIGANICS 1 81071 808 (0108/DEO2) OHGANICS \$12% and Tare (0508/0502) select bre 7218 BIOU BOS (IUS 108/0E/05) BUNOSEB HOL × X X ANALYTES (METHOD) X × × یز × X NO. OF CONTAINERS 3 J 7 RECEIVED BY: (SIGNATURE) water water water water water Water Water water Water Water Matrix water water water Depth 1 ١ Begin Depth l l Project Manager: Todd Wiedemeier (Denver, CO) THE Project Name/Location: Beale AFB, UST Site 3/11/96/1400 3/12/96/1420 Site Manager: Michael Phelps (Alameda, CA) 1345 Time 3(12(94) 1430 DATE 3/12/96 3/12/96 Sampler(s): (Initials and Signature(s)) 3/12/96 Date RELINQUISHED BY: (SIGNATURE) Project No.: 722450.14020 MAR Made BE-MP46-B BE-MP08-01 BE-MP41-0 BE-MP45-Ø1 BE-MP414-0 BE-MPII- DI Sample ID

'stribution. Original accompanies shipment; photocopy kept by samplers; copy FAXed to Parsons ES Site Manager by laboratory upon receipt of samples.

4. TMBs: 1,2,3-trimethylbenzene, 1,2,4-trimethylbenzene, and 1,3,5-

Secondary column confirmation required.
 Quantify all carbon ranges: diesel, jet fuel, oll/grease.
 Delonized water extraction

REMARKS/COMMENTS:

TIME

REGEIVED FOR LAB BY: (SIGNATURE) DATE

TIME

DATE

RELINGUISHED BY: (SIGNATURE)

trimethylbenzene.

COOLER RECEIPT CHECKLIST



Rev. 1 4/95

Login	#: 124797 Date Received: 3/13/96 Number of Coolers:	/
Client	: Parons Ene Project: Beale	
A.	Preliminary Examination Phase Date Opened: 3/13/96 By (print): Leveling Settles (sign)	lu.
1.	Did cooler come with a shipping slip (airbill, etc.)? If YES, enter carrier name and airbill number: Persex 11540927	YES NO
2.	Were custody seals on outside of cooler?	(YES) NO
~	How many and where? Seal date: Seal name:	
3.	Were custody seals unbroken and intact at the date and time of arrival?	.(YES) NO
4.	Were custody papers dry and intact when received?	. NO
5.	Were custody papers filled out properly (ink, signed, etc.)?	ўт ў NO
6.	Did you sign the custody papers in the appropriate place?	. YE'S NO
7.	Was project identifiable from custody papers?	YES NO
0	If YES, enter project name at the top of this form.	
8.	If required, was sufficient ice used?	(YÉS) NO
	Type of ice: Yeal Temperature: 4°C	
B.	Login Phase Date Logged In: 3/13/96 By (print): Cynthi Seller (sign) Contlud	Lale
1.	Describe type of packing in cooler: pupu beable unas	- Seniely
2.	Did all bottles arrive unbroken?	(YES NO
3.	Were labels in good condition and complete (ID, date, time, signature, etc.)?	(YES) NO
4.	Did bottle labels agree with custody papers?	NO NO
5.	Were appropriate containers used for the tests indicated?	VES NO
6.	Were correct preservatives added to samples?	(YES NO
7.	Was sufficient amount of sample sent for tests indicated?	YESINO
3.	Were bubbles absent in VOA samples? If NO, list sample Ids below	YESINO
9.	Was the client contacted concerning this sample delivery?	YES NO
	If YES, give details below.	125 110
	Who was called? By whom? Date:	
Additio	nal Comments:	
	·	
		
	·	·
Filename: i	F:\qc\forms\cooler.wpd	1 4/95



Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 94710, Phone (510) 486-0900

ANALYTICAL REPORT

Prepared for:

Parsons Engineering Science, Inc. 1301 Marina Village Parkway Suite 200 Alameda, CA 94501

Date: 02-APR-96

Lab Job Number: 124833

Project ID: 722450.14020 Location: Beale AFB

Reviewed by:

Reviewed by:

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MEMUKANDUM TO FILE

211 100000

JOB NO. 722450, 14020

	FILE DESIGNATION Beale AFB - UST 5:te
	DATE 6/14/96 TIME
PHONE CALL TO	PHONE NO
CONFERENCE WITH	
PLACE	
SUBJECT Resampled monitoring points MP. ambient blank (MP-47) due to	
Due to missed holding times	for samples collected
on 3/13/96, The wells were re:	sampled on 5/14/96 and
5/15/96.	
	·
	Mand Man -



Laboratory Number: 124833

Client: Parsons Engineering Science

Project#: 722450.14020 Location: Beale AFB Sample Date: 03/13/96 Receipt Date: 03/14/96

CASE NARRATIVE

This hardcopy data package contains sample results and batch QC for four water samples and one trip blank received from the above referenced project on March 14, 1996. Two of the VOA vials for sample BE-MP13-01 were broken upon receipt. No TPH/diesel results will be available for sample BE-MP12-01 as no liter amber bottle was submitted. A trip blank was submitted with these samples but was not documented on the chain-of-custody. Per the client request, the trip blank was analyzed for TPH/gasoline and BTXE/TMBs.

TPH/Extractable: No analytical problems were encountered.

TPH/Purgeable: Due to instrument problems the TPH/gasoline holding times were missed by one day. The samples affected are BE-MP10-01 (C&T# 124833-001), BE-MP12-01 (C&T# 124833-002), BE-MP13-01 (C&T# 124833-003) and BE-MP47-01 (C&T# 124833-004). Bromobenzene failed surrogate recovery limits for sample BE-MP10-01 (C&T# 124833-001), possibly due to coelution of sample hydrocarbons with this surrogate.

BTXE (EPA 8020): Due to instrument problems the BTXE/TMB holding times were missed by one day. The samples affected are BE-MP10-01 (C&T# 124833-001), BE-MP12-01 (C&T# 124833-002), BE-MP13-01 (C&T# 124833-003) and BE-MP47-01 (C&T# 124833-004).



Chromatograms are included in this report for any samples containing quantifiable concentrations of extractable or purgeable hydrocarbons, along with example chromatograms for each type of fuel.

Data Qualifier Flags

C: The "C" qualifier flag indicates that the target compound reported was confirmed by a second analytical column, but that the concentration quantitated on the second column differed from the reported concentration by more than a factor of two. This often happens when the fuel pattern present in the samples does not resemble gasoline.

DO: The surrogate recovery could not be calculated due to the dilution required for sample analysis.

L: The "L" qualifier flag indicates that the sample chromatogram resembles a fuel pattern, but that the hydrocarbons present are lighter than those seen in a standard of the reported fuel.

H: The "H" qualifier flag indicates that the sample chromatogram resembles a fuel pattern, but that the hydrocarbons present are heavier than those seen in a standard of the reported fuel.

Y: The "Y" qualifier flag is used to indicate that the sample chromatogram does not resemble the standard for the reported fuel range. This flag is often used in combination with the flags "L" and/or "H".

Z: The "Z" qualifier flag indicates that the sample chromatogram does not resemble a fuel pattern, but consists of individual, strongly differentiated peaks.



Page 1 of 1

TEH-Tot Ext Hydrocarbons

client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Location: Beale AFB Prep Method: EPA 3520

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124833-001 BE-MP10-01	26564	03/13/96	03/21/96	03/25/96	
124833-003 BE-MP13-01	26564	03/13/96	03/21/96	03/25/96	

Analyte Diln Fac:	Units	124833- 1	001	124833- 1	003	
JP5	ug/L	1500	YH	2200	YH	
Diesel Range	ug/L	4300	YLH	3500	YHL	
Motor Oil Range	ug/L	2200	YL	1600	YL	
Surrogate					-	
Hexacosane	%REC	74		87		

Sample exhibits fuel pattern which does not resemble standard

H: Heavier hydrocarbons than indicated standard

L: Lighter hydrocarbons than indicated standard



BATCH QC REPORT

Page 1 of '

TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M

Project#: 722450.14020

EPA 3520

Location: Beale AFB

Prep Method:

METHOD BLANK

Matrix: Water Batch#: 26564 Units: ug/L

03/21/96 Prep Date: Analysis Date: 03/25/96

Diln Fac: 1

MB Lab ID: QC17632

Analyte	Result	
JP5	<50	
Diesel Range	<50	
Motor Oil Range	<300	
Surrogate	%Rec	Recovery Limits
Hexacosane	99	60-140



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Method:

EPA 3520 Location: Beale AFB

BLANK SPIKE/BLANK SPIKE DUPLICATE

Matrix: Water Prep Date: 03/21/96 03/25/96

26564 Batch#: Analysis Date: Units: ug/L Diln Fac: 1

BS Lab ID: QC17633

Analyte	Spike Added	BS	%Rec #	Limits
Diesel Range	2475 22	65	92	60-140
Surrogate	%Rec	Limits		
Hexacosane	90	60-140	W. S. W. W. W. W. W. W. W. W. W. W. W. W. W.	

BSD Lab ID: QC17634

Analyte	Spike Added	BSD	%Rec #	Limits	RPD #	Limit
iesel Range	2475	2355	95	60-140	4	<35
Surrogate	%Rec	Limit	ts			
Hexacosane	88	60-14	40			

[#] Column to be used to flag recovery and RPD values with an asterisk

RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits

CHAINTET TOD

Sample Name : S,124833-001,26564 FileName : C:\GC15\CHB\084B016.RAW

Method : BTEH.MTH

Start Time : 0.01 min Scale Factor: 0.0

End Time : 31.91 min

Plot Offset: 41 mV

Sample #: 500:2.5

Date: 3/27/96 10:51 AM

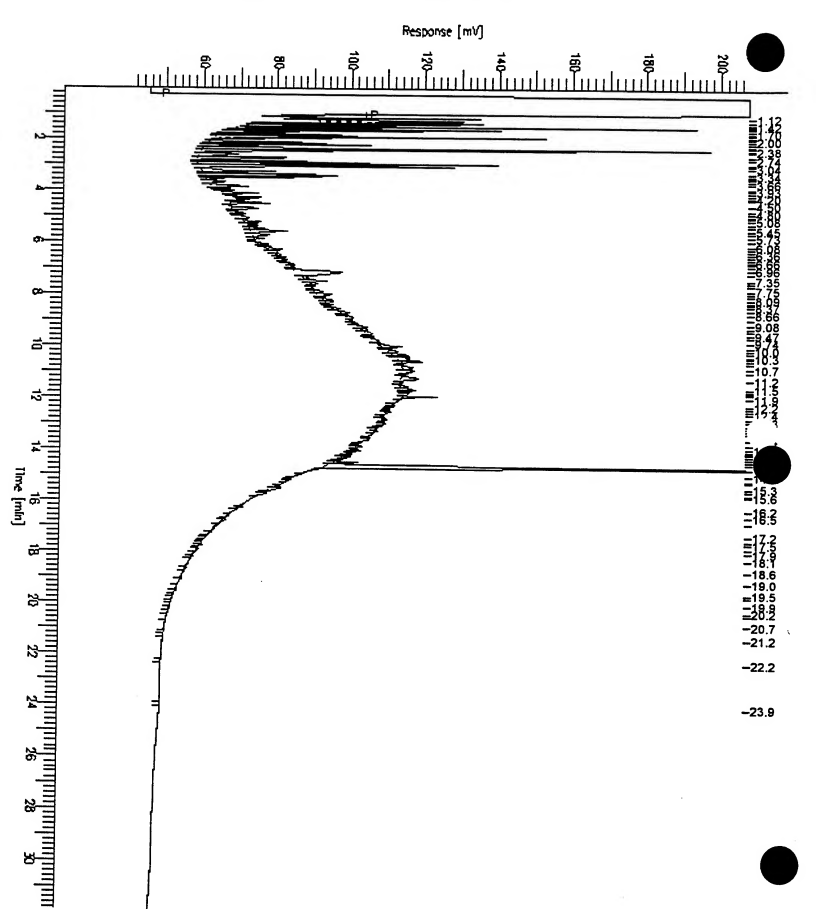
09:09 AM

Time of Injection: 3/25/96 Low Point: 40.83 mV

High Point : 207.76 mV

Page 1 of 1

Plot Scale: 166.9 mV



GC15 Channel A TEH

Sample Name : S,124833-003,26564

FileName : C:\GC15\CHB\084B017.RAW

Method : BTEH.MTH

Start Time : 0.01 min S · Factor: 0.0

End Time : 31.91 min

Plot Offset: 38 mV

Sample #: 500:2.5

ample #: 500:2.5

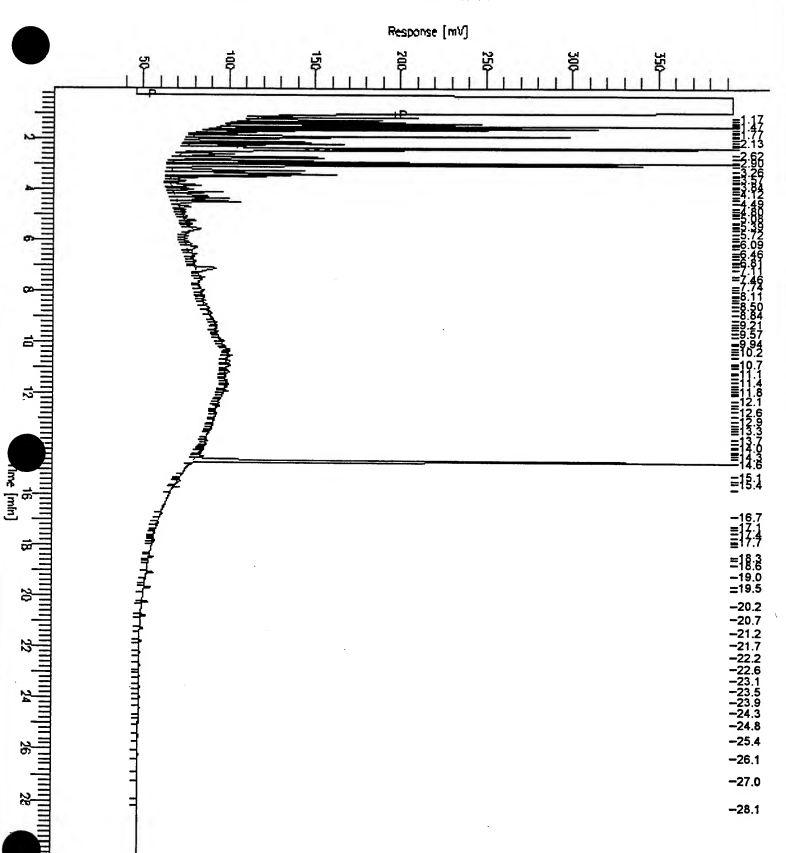
Date: 3/27/96 10:53 AM

Time of Injection: 3/25/96 09:53 AM

Low Point: 37.93 mV High Point: 393.26 mV

Page 1 of 1

Plot Scale: 355.3 mV



JP-5

GC15 Channel A TEH

Sample Name : CCV,96WS2069,JP5
FileName : C:\GC15\CHB\084B023.RAW

Method : STEH.MTH

Start Time : 0.01 min Scale Factor: 0.0

End Time : 19.80 min Plot Offset: 42 mV

Sample #: 250MG/L

Date: 3/26/96 12:54 PM

Time of Injection: 3/25/96 01:29 PM

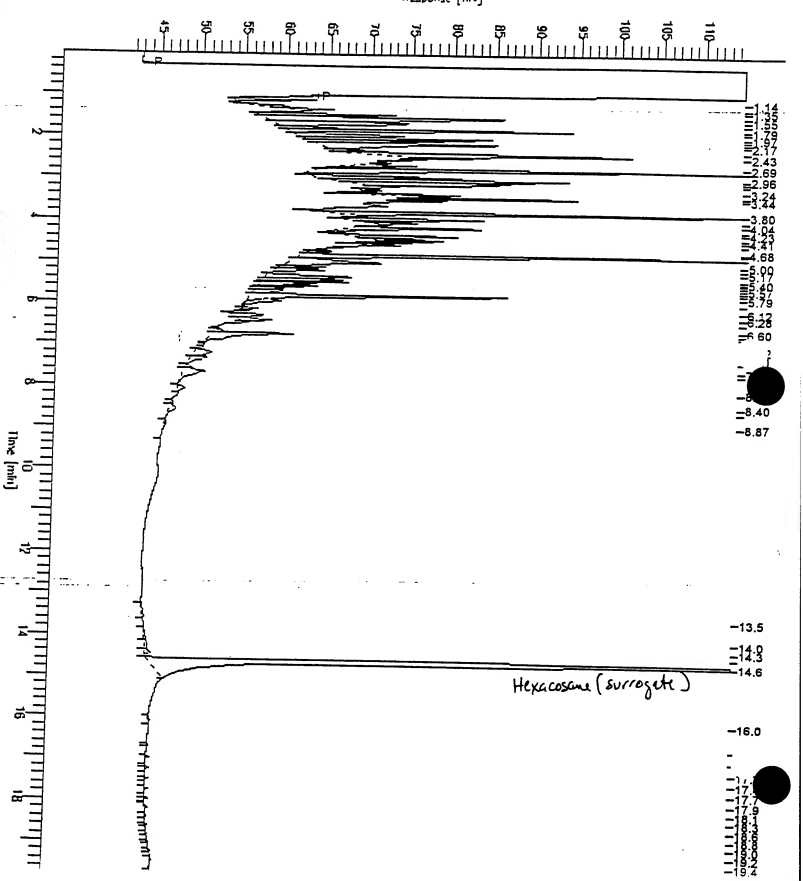
Low Point : 41.54 mV

High Point : 114.59 mV

Page 1 of 1

Plot Scale: 73.1 mV





Sample Name : CCV, 96WS1948, DSL

FileName : C:\GC15\CHB\084B039.RAW

Method : BTEH.MTH

Start Time : 0.01 min Scale Factor: 0.0

End Time : 31.91 min

Plot Offset: 40 mV

Sample #: 500MG/L

Date: 3/26/96 12:58 PM

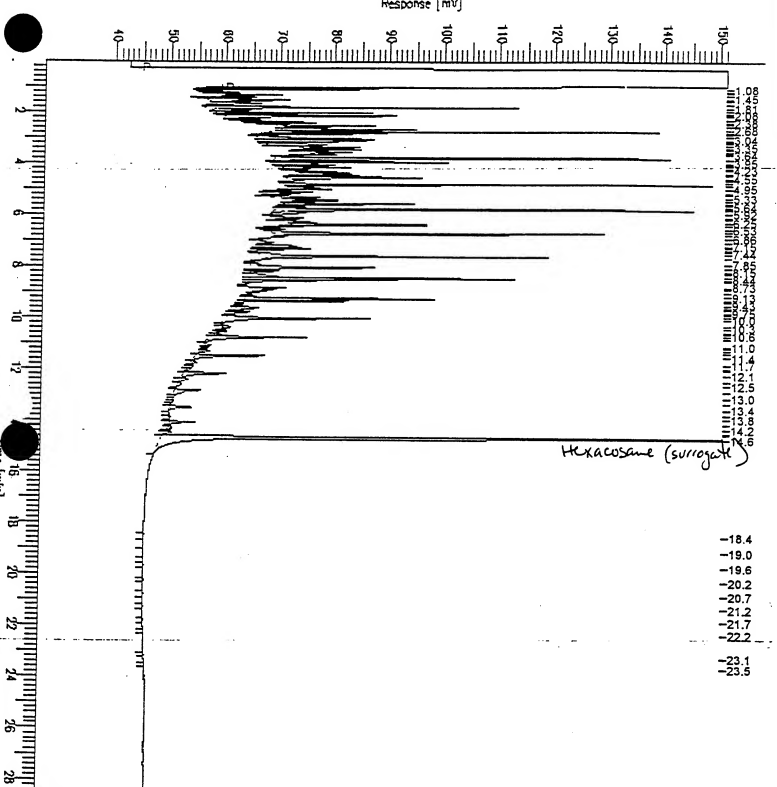
Time of Injection: 3/26/96 12:64 AM Low Point : 39.68 mV

High Point : 151.11 mV

Page 1 of 1

Plot Scale: 111.4 mV





motor

Sample Name : CCV,96WS2112,MO
FileName : C:\GC15\CHB\084B042.RAW

Hethod : BTEH.MTH

Start Time : 0.01 min Scale Factor: 0.0

End Time : 31.91 min

Plot Offset: 25 mV

Sample #: 500MG/L

Date: 3/26/96 12:55 PM

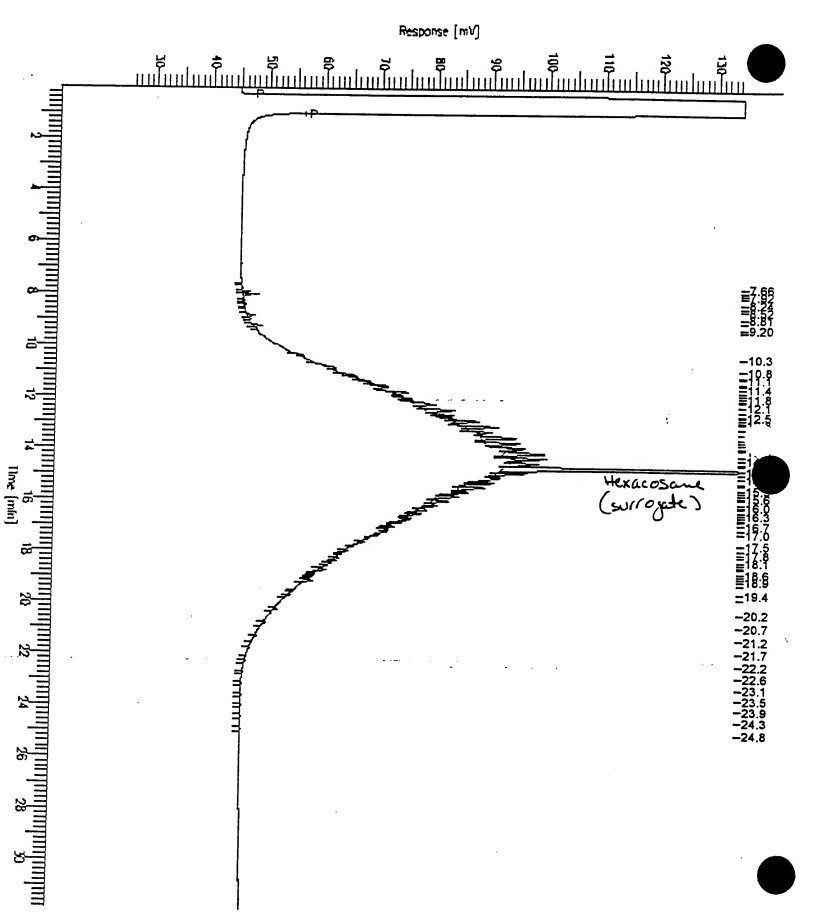
Time of Injection: 3/26/96 02:17 AM

Low Point : 25.07 mV

High Point : 134.45 mV

Page 1 of 1

Plot Scale: 109.4 mV





TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 5030

Locat	ion:	Beale	AFB
-------	------	-------	-----

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124833-001 BE-MP10-01	26672	03/13/96	03/28/96	03/28/96	
124833-002 BE-MP12-01	26672	03/13/96	03/28/96	03/28/96	
124833-003 BE-MP13-01	26672	03/13/96	03/28/96	03/28/96	
124833-004 BE-MP47-01	26672	03/13/96	03/28/96	03/28/96	

Analyte Diln Fac:	Units	124833 1	-001	124833-002 1	124833-003 10	124833-004 1
Gasoline	ug/L	1600)	<50	5200	<50
Surrogate						
Trifluorotoluene	%REC	104		99	99	100
Bromobenzene	%REC	125	*	105	105	98

Values outside of QC limits



TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015)

Project#: 722450.14020 Location: Beale AFB Prep Method: EPA 5030

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124833-005	TRIP BLANK	26672	03/13/96	03/27/96	03/27/96	

Analyte Diln Fac:	Units	124833-005 1		
Gasoline	ug/L	<50		
Surrogate				
Trifluorotoluene	%REC	99	 	
Bromobenzene	%REC	95		



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

METHOD BLANK

Matrix: Water

Prep Date: 03/27/96

Batch#: 26672 Units: ug/L Analysis Date: 03/27/96

Diln Fac: 1

MB Lab ID: QC18055

Result	
<50	
%Rec	Recovery Limits
95	69-120
87	70-122
	<50 %Rec



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Parsons Engineering Science, Inc. Client:

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26672 Units:

ug/L Diln Fac: 1

03/27/96 Prep Date: 03/27/96 Analysis Date:

LCS Lab ID: QC18056

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	1843	2006	92	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	103 101	69-120 70-122		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits

BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Method: EPA 5030

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

 Field ID:
 ZZZZZZZ
 Sample Date:
 03/12/96

 Lab ID:
 124797-006
 Received Date:
 03/13/96

 Matrix:
 Water
 Prep Date:
 03/27/96

 Batch#:
 26672
 Analysis Date:
 03/27/96

Units: ug/L Diln Fac: 1

MS Lab ID: QC18058

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Gasoline	2000	<50.00	2047	102	75-125
Surrogate	Surrogate %Rec				
Trifluorotoluene Bromobenzene	104 110	69-120 70-122			

MSD Lab ID: QC18059

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Gasoline	2000	2135	107	75-125	4	<20
Surrogate	%Rec	Limits				
Trifluorotoluene Bromobenzene	104 114	69-120 70-122				

[#] Column to be used to flag recovery and RPD values with an asterisk

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits RPD: 0 out of 1 outside limits

: G:\GC05\087H026.raw

Start Time : 0.00 min

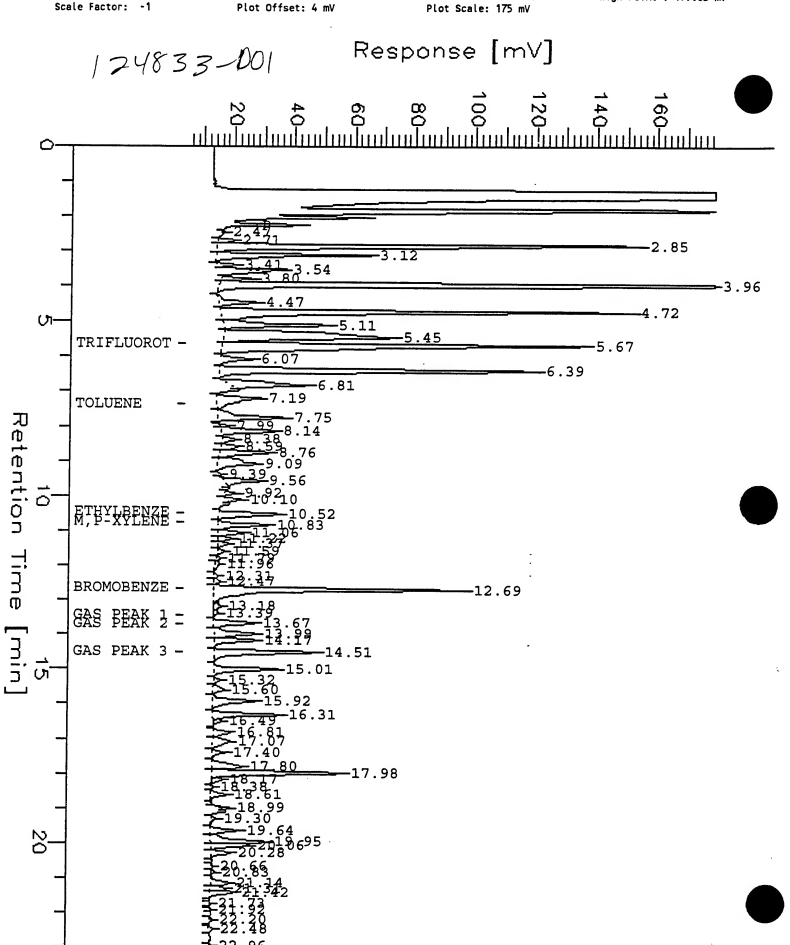
23.35

End Time : 23.42 min

Date: 3/28/96 5:51 AM

Low Point : 4.03 mV

Page 1 of 1 High Point: 179.03 mV



FileName : G:\GC05\087H025.raw Start Time : 0.00 min Er

End Time : 23.42 min Plot Offset: 4 mV

-23.36

Date: 3/28/96 5:13 AM Low Point : 4.07 mV

Plot Scale: 175 mV

Page 1 of 1 High Point : 179.07 mV

Scale Factor: -1 Response [mV] 124833-3 2.85 .13 23.41 -3.96 4.46 4.73 5.11 -5.45 UT-TRIFLUOROT -5.68 6.08 6.39 .30 = 7.75 818 577 -6.81 TOLUENE ZJ E on Time [min] BROMOBENZE --12.69 GAS PEAK 1 = 13.67 13:95 14:17 ---14.51 GAS PEAK 3 -15.00 (A) 5.59 5.92 -18.99 19.29 -19.64 20.28 20.64

TVH2 - GC-04 RTX-1

C+T Gasoline Std.

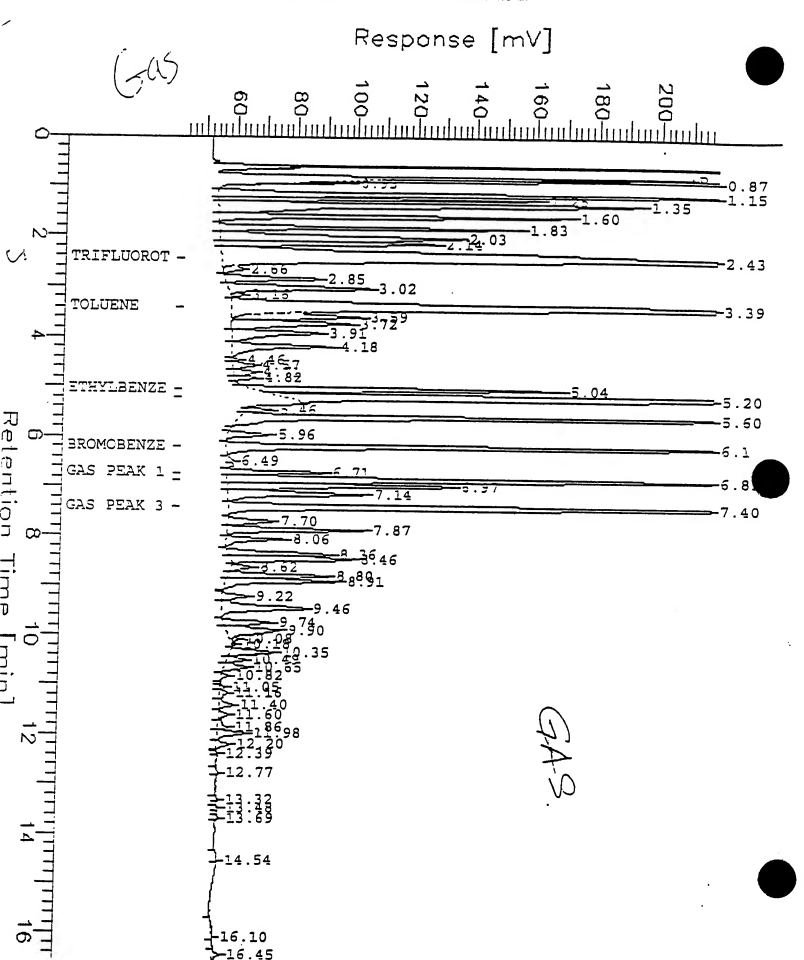
FileName : G:\GC04\135J001.raw

Start Time : 0.00 min Scale Factor: -1

End Time : 17.00 min Plot Offset: 43 mV Date : 5/15/95 9:53 AM Low Point : 42.92 mV

Plot Scale: 175 mV

Page 1 of 1 High Point : 217.92 mV



TVH2 - GC-04 RTX-1 FileName : G:\GC04\B\026J011.raw Date : 2/1/95 3:57 PM Low Point : -4.15 mV Start Time : 0.00 min End Time : 17.00 min High Point: 170.85 mV Plot Offset: -4 mV Scale Factor: -1 Plot Scale: 175 mV Response [mV] 0.93 11. (190 -1.25 -1.74 <u>-2.01</u> 33 -2.14 TRIFLUOROT -TOLUENE -3.62 $\frac{-3.81}{4.02}$ 4.38 4.58 5.06 4.96 .28 M, H X DEENE = -5.44 5.76.57 5.9785 6.20 BROMOBENZE -GAS PEAK 1 = GAS PEAK 3 -7.65 \otimes -7.93 9.70 9.55 -10.02

Retention

Time

min

Page 1 of 1

-1.46

2.66 -3.06



BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124833-001 BE-MP10-01	26672	03/13/96	03/28/96	03/28/96	
124833-002 BE-MP12-01	26672	03/13/96	03/28/96	03/28/96	
124833-003 BE-MP13-01	26672	03/13/96	03/28/96	03/28/96	
124833-004 BE-MP47-01	26672	03/13/96	03/28/96	03/28/96	

Analyte till Diln Fac:	Jnits	124833-001 1	124833-002 1	124833-003 10	124833-004
	- 14	-	-		
Benzene	ug/L	<0.5	<0.5	<5	<0.5
Toluene	ug/L	<0.5	<0.5	<5	<0.5
Ethylbenzene	ug/L	18	<0.5	5.5C	<0.5
m,p-Xylenes	ug/L	14	<0.5	12	<0.5
o-Xylene	ug/L	<0.5	<0.5	<5	<0.5
1,2,3-Trimethylbenzene	ug/L	14	<0.5	37	<0.5
1,2,4-Trimethylbenzene	ug/L	20	<0.5	54	<0.5
1,3,5-Trimethylbenzene	ug/L	16 C	<0.5	31 C	<0.5
Surrogate					
Trifluorotoluene	%REC	102	95	98	96
Bromobenzene	%REC	103	89	93	91

C: Presence of this compound confirmed by second column, however, the confirmation concentration differed from the reported result by more than a factor of two



BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB

Analysis Method: EPA 8020 Prep Method: EPA 5030

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124833-005	TRIP BLANK	26672	03/13/96	03/27/96	03/27/96	

Analyte	Jnits	124833-005	
Diln Fac:		1	
Benzene	ug/L	<0.5	
Toluene	ug/L	<0.5	
Ethylbenzene	ug/L	<0.5	
m,p-Xylenes	ug/L	<0.5	
o-Xylene	ug/L	<0.5	
1,2,3-Trimethylbenzene	e ug/L	<0.5	
1,2,4-Trimethylbenzene	e ug/L	<0.5	
1,3,5-Trimethylbenzene	e ug/L	<0.5	
Surrogate			
Trifluorotoluene	%REC	99	
Bromobenzene	%REC	89	



BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

METHOD BLANK

Matrix: Water Batch#: 26672 Units: ug/L Diln Fac: 1

Prep Date: 03/27/96 Analysis Date: 03/27/96

MB Lab ID: QC18055

Analyte	Result	
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene	94	58-130
Bromobenzene .	81	62-131



BATCH QC REPORT

Page 1 of 1

BTXE

lient: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Prep Date: 03/27/96

Matrix: Water Batch#: 26672 Analysis Date: 03/27/96

Units: ug/L Diln Fac: 1

LCS Lab ID: QC18057

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	16.8	20	84	80-120
Toluene	17.7	20	89	80-120
Ethylbenzene	16.2	20	81	80-120
m,p-Xylenes	36.8	40	92	80-120
o-Xylene	17.5	20	88	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	96	58-130	·	
Bromobenzene	81	62-131		

Column to be used to flag recovery and RPD values with an asterisk Values outside of QC limits

ike Recovery: 0 out of 5 outside limits

LOGIN CHANGE FORM CURTIS & TOMPKINS, LTD. BERKELEY

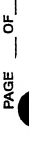
3/18/96 Inilials: The Client Request: By: Michael Fheles Date/Time:
Login Review Data Review Reason for change:

Duedale	3/27									
Ánalysis	TUHBAMB	/ /		ī						
Add/Caucol	430 Add									
Matrix	H20									٠
Cllent ID	TRIP RIANE)								
Provious Lab ID										
Curront Lab ID	124833-5	•								



1301 Marina Village Pkwy, Suite 200 Alameda, California 94501 Phone: (510) 769-0100 FAX: (510) 769-9244





Project No.: 722450.14020	20								ORGANICS	1111	INORGANICS	OTHER	
Project Name/Location: Beale AFB, UST Site	Beale AFB,	UST Site	-				(-	1/2/				
Project Manager: Todd Wiedemeier (Denver, CO)	Wiedemeier	(Denver,	(00				POH	1 0	\$910	////	//	///	
Site Manager: Michael Phelps (Alameda, CA)	helps (Alam	eda, CA)					LAW		903 (O	///		1// (9,0	
Sampler(s): (Initials and Signature(s))	Signature(s)	7					S3 (WS108	\$100/0	108/08/05				•
N N) .	14 Many	1 /			10.0FC	JANA BAOEOZ) ONT BAOEOZ) ONT BAOEOZ)	\	202) 28MT 200V DE (0758/02?			Note: All analyses and holding times	
Sample ID	Date	TIMe	Begin Depth	End Depth	Matrix	HAY	MARIOR DES	STENOSONEH STENOSONEH	E) SO			According to AFCEE handbook requirements Remarks	a i
RE-MPIG-01	3/1396	13960935)	1	water	4 1/2	Z	2					
BE-MP12-01	3/13/96	13/96/1015		1		メメ	\ \ \ \	×				No Liter Awler for TEH ON WIL	
	3/13/96	1,155	1	1	water X	×	7	7				2 vons acriged	
	3/13/46		١	١	water	3 次		7				mjosa	
					water		E 27						
Triphak					water	*		米				* Eur Der MP. 5/	18/
					water								1700
					water								
		N 40 1			water								
					water				7 10				
					water		9.73						
					water								
					water								
RELINQUISHED BY: (SIGNATURE)	ATURE) DATE			EIVED BY	RECEIVED BY: (SIGNATURE)	(9)	NOTES: 1. Seco 2. Quan 3. Delor	40TES: 1. Secondary column confirm 2. Quantity all carbon ranges: 3. Delonized water extraction	IOTES: 1. Secondary column confirmation required 2. Cuantify all carbon ranges: diesel, jet fue 3. Deionized water extraction	10 TES: 1. Secondary column confirmation required. 2. Quantify all carbon ranges: diesel, jet fuel, oll/grease. 3. Deionized water extraction		4. TMBs: 1,2,3-timethylbenzene, 1,2,4 trimethylbenzene, and 1,3,5- trimethylbenzene.	*
RELINGUISHED BY (SIGNATURE)	ATURE) DATE		TIME REC	EIVED FO	RECEIVED FOR LAB BY: (\$	(SIGNATURE)	<u> </u>	TIME	REMARKS	REMARKS/COMMENTS:			
The form	7/	13/96 1400	90		Imaa M	(arrist	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	200					
							1						1

Distribution: Original accompanies shipment; photocopy kept by samplers; copy FAXed to Parsons ES Site Manager by laboratory upon receipt of samples.

COOLER RECEIPT CHECKLIST Curtis & Tompkins, Lid.



Login	
Client	: Varsens E.S. Project: Beale AFR
A.	Preliminary Examination Phase Date Opened: 3 14 96 By (print): T. Marcis (Man) T. Marcis T. Marcis (Man)
1.	Did cooler come with a shipping slip (airbill, etc.)? If YES, enter carrier name and airbill number: F5D Sv 115 409 215 6
2.	Were custody seals on outside of cooler? How many and where? Front / 18ac Seal date: 5/13 Seal name: H. S. Chi 1
3.	Were custody seals unbroken and intact at the date and time of arrival?
4.	Were custody papers dry and intact when received?
5.	Were custody papers filled out properly (ink, signed, etc.)?
6.	Did you sign the custody papers in the appropriate place?
7.	Was project identifiable from custody papers? NO
	If YES, enter project name at the top of this form.
8.	If required, was sufficient ice used?
	Type of ice: WET Temperature: 6°C
B.	Login Phase Date Logged In: By (print): JWI (sign) 4-6-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-
1.	Describe type of packing in cooler:
<u>2</u>)	Did all bottles arrive unbroken? YES NO
3.	Were labels in good condition and complete (ID, date, time, signature, etc.)?. YBS NO
€	Did bottle labels agree with custody papers? YES
5.	Were appropriate containers used for the tests indicated?
5.	Were correct preservatives added to samples?
7.	Was sufficient amount of sample sent for tests indicated? YES NO
3.	Were bubbles absent in VOA samples? If NO, list sample Ids below
₽.	Was the client contacted concerning this sample delivery? YES NO If YES, give details below.
	Who was called? By whom? Date:
2) * No Trip	nal Comments: only I VOA VICT red intent for BE-MPOIZ-DI Liter Auba submitted for TEH sugar BEMP12-01 -don 3/15 bank received based on these for 3/15
lename: f	:\qc\forms\cooler.wpd



Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 94710, Phone (510) 486-0900

ANALYTICAL REPORT

Prepared for:

Parsons Engineering Science, Inc. 1301 Marina Village Parkway Suite 200 Alameda, CA 94501

Date: 02-APR-96

Lab Job Number: 124895

Project ID: 722450.14020 Location: Beale AFB

Reviewed by:

Reviewed by:

This package may be reproduced only in its entirety.



Laboratory Number: 124895

Client: Parsons Engineering Science

Project#: 722450.14020 Location: Beale AFB Sample Date: 03/20/96 Receipt Date: 03/20/96

CASE NARRATIVE

This hardcopy data package contains sample results and batch QC for one trip blank, one water sample and one composite soil sample received from the above referenced project on March 20, 1996. All samples were received cold and intact. The trip blank was not documented on the chain-of-custody. Per the client request, the trip blank was analyzed for TPH/gasoline and BTXE/TMBs.

TPH/Extractable: No analytical problems were encountered.

TPH/Purgeable: No analytical problems were encountered.

BTXE (EPA 8020): No analytical problems were encountered.



Page 1 of 1

TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 3520

Location: Beale AFB

Sample # (Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124895-001 B	BE-IDW-01	26564	03/20/96	03/21/96	03/25/96	

Analyte Diln Fac:	Units	124895 - 1	-001		
JP5	ug/L	780	YH		
Diesel Range	ug/L	1100	YLH		
Motor Oil Range	ug/L	460	YL		
Surrogate					
Hexacosane	%REC	88		 	

Sample exhibits fuel pattern which does not resemble standard at Heavier hydrocarbons than indicated standard

L: Lighter hydrocarbons than indicated standard



TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: LUFT

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124895-007	COMP IDW-(2,3,4,5,6)	26567	03/20/96	03/21/96	03/27/96	17%

Analyte Diln Fac:	Units	124895-007 1	
JP5 Diesel Range Motor Oil Range	mg/Kg mg/Kg mg/Kg	3.1YH 17 YH 110 YH	
Surrogate			
Hexacosane	%REC	109	

Y: Sample exhibits fuel pattern which does not resemble standard

H: Heavier hydrocarbons than indicated standard



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 3520

Location: Beale AFB

METHOD BLANK

Prep Date: 03/21/96 Analysis Date: 03/25/96

Matrix: Water Batch#: 26564 Units: ug/L Diln Fac: 1

MB Lab ID: QC17632

Analyte	Result	
JP5	<50	
Diesel Range	<50	
Motor Oil Range	<300	
Surrogate	%Rec	Recovery Limits
Hexacosane	99	60-140



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Parsons Engineering Science, Inc. Client:

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: LUFT

METHOD BLANK

Matrix: Soil 26567 Batch#:

Prep Date: Analysis Date:

03/21/96 03/25/96

Units: mg/Kg Diln Fac: 1

MB Lab ID: QC17643

Analyte	Result	
JP5	<1.0	
Diesel Range	<1.0	
Motor Oil Range	<5.0	
Surrogate	%Rec	Recovery Limits
Hexacosane	139	60-140



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method:

EPA 3520

Location: Beale AFB

BLANK SPIKE/BLANK SPIKE DUPLICATE

03/21/96

Matrix: Water 26564 Batch#: Units: ug/L

Diln Fac: 1

Prep Date: Analysis Date: 03/25/96

BS Lab ID: QC17633

Analyte	Spike Added	BS	%Rec #	Limits
Diesel Range	2475 2	265	92	60-140
Surrogate	%Rec	Limits		
Hexacosane	90	60-140	-	

BSD Lab ID: QC17634

Analyte	Spike Added	BSD	%Rec #	Limits	RPD #	Limit
iesel Range	2475	2355	95	60-140	4	<35
Surrogate	%Rec	Limits	5			
Hexacosane	88	60-140)			12 00.0

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Parsons Engineering Science, Inc. Client:

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method:

Location: Beale AFB

LABORATORY CONTROL SAMPLE

03/21/96 Prep Date:

Matrix: Soil Batch#: 26567 Analysis Date: 03/25/96

Units: mg/Kg Diln Fac: 1

LCS Lab ID: QC17644

Analyte	Result	Spike Added	%Rec #	Limits
Diesel Range	55.4	49.5	112	60-140
Surrogate	%Rec	Limits		
Hexacosane	125	60-140		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits

BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: LUFT

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: COMP IDW-(2,3,4,5,6)

Sample Date:

03/20/96

Lab ID: 124895-007 Matrix: Soil

Received Date:

03/20/96 03/21/96

Prep Date:

03/25/96

Batch#: 26567

Analysis Date:

Units: mg/Kg dry weight

17%

Diln Fac: 1

Moisture:

MS Lab ID: QC17645

Analyte	Spike Added	Sample	MS	%Rec #	Limits	
Diesel Range	59.64	16.87	62.89	77	60-140	
Surrogate	%Rec	%Rec Limits				
Hexacosane	132	60-140				

4SD Lab ID: QC17646

nalyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Diesel Range	59.64	57.47	68	60-140	13	<30
Surrogate	%Rec	Limits				
Hexacosane	120	60-14	0			

[#] Column to be used to flag recovery and RPD values with an asterisk

RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits

cnannel GLID A TEH

Sample Name : S,124895-001,26564 FileName : C:\GC15\CHB\084B015.RAW

: BTEH.MTH Method

Start Time : 0.01 min Scale Factor: 0.0

End Time : 31.91 min - Plot Offset: 43 mV

Sample #: 500:2.5

Date: 3/26/96 02:48 PM

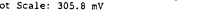
Time of Injection: 3/25/96 08:24 AM

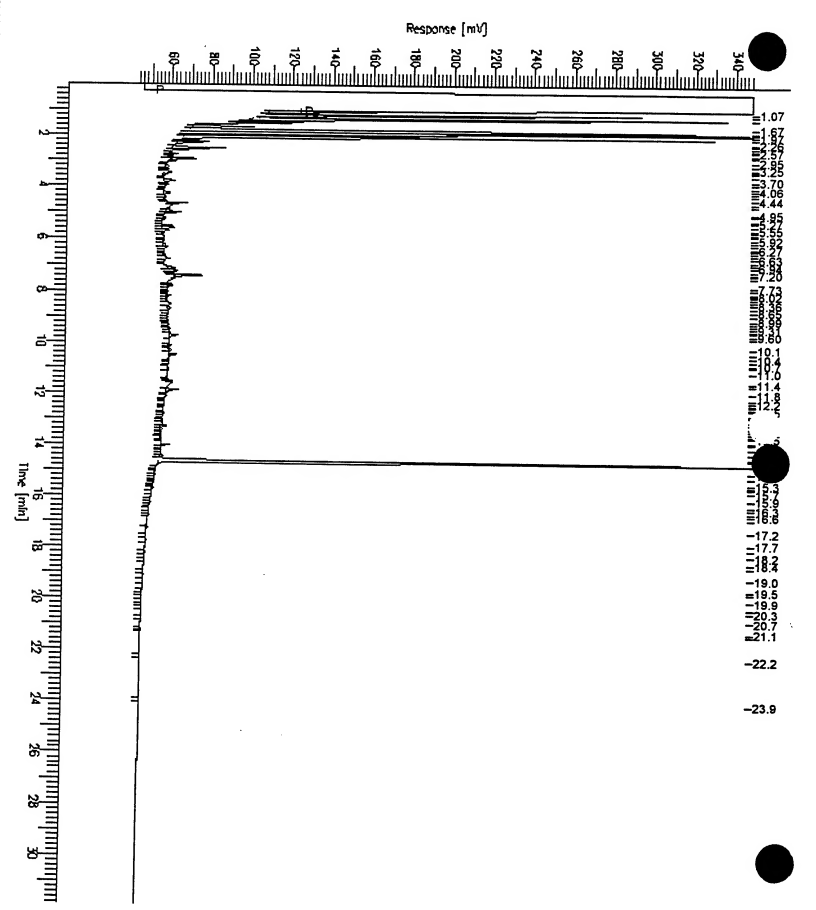
Low Point: 42.54 mV

Plot Scale: 305.8 mV

High Point : 348.33 mV

Page 1 of 1





TEH Chromatogram-GC13 CH A

Sample Name: 124895-007,50:5

: g:\gc13\cha\087A021.raw FileName

: 13DUAL32.ins Method

Start Time : 0.00 min Scale Factor: -1

End Time : 31.92 min

Plot Offset: 34 mV

Sample #: 26567 Date : 3/27/96 09:51 PM

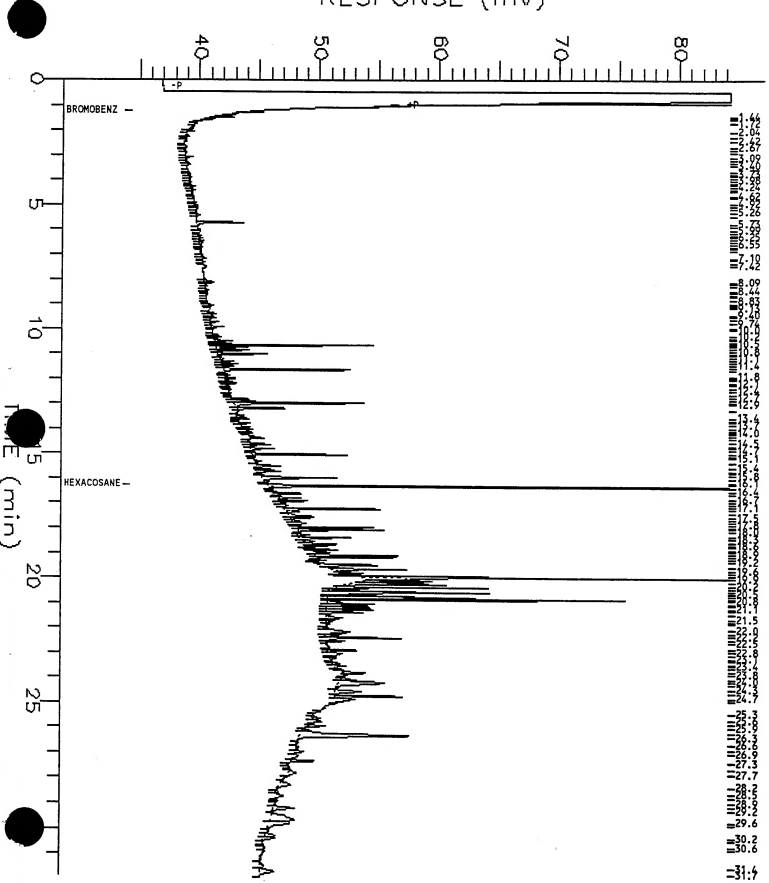
Time of Injection: 3/27/96 09:17 PM Low Point: 34.32 mV High Po

Plot Scale: 50 mV

High Point: 84.32 mV

Page 1 of 1





Sample Name : CCV, 96WS1948, DSL Sample #: 500MG/L Page 1 of 1 Date: 3/26/96 12:58 PM FileName C:\GC15\CHB\084B039.RAW Method BTEH. MTH Time of Injection: 3/26/96 12:04 AM Start Time : 0.01 min End Time : 31.91 min Low Point : 39.68 mV High Point : 151.11 mV Scale Factor: 0.0 Plot Offset: 40 mV Plot Scale: 111.4 mV usel Response [mV]

-18.4 -19.0

-19.6 -20.2 -20.7 -21.2 -21.7 -22.2

-23.1 -23.5

Sample Name: CCV, 96WS2112, MO

FileName : C:\GC15\CHB\084B042.RAW

Method : BTEH.MTH

: 0.01 min Start Time Scale Factor:

End Time : 31.91 min

Plot Offset: 25 mV

Sample #: 500MG/L

Date: 3/26/96 12:55 PM

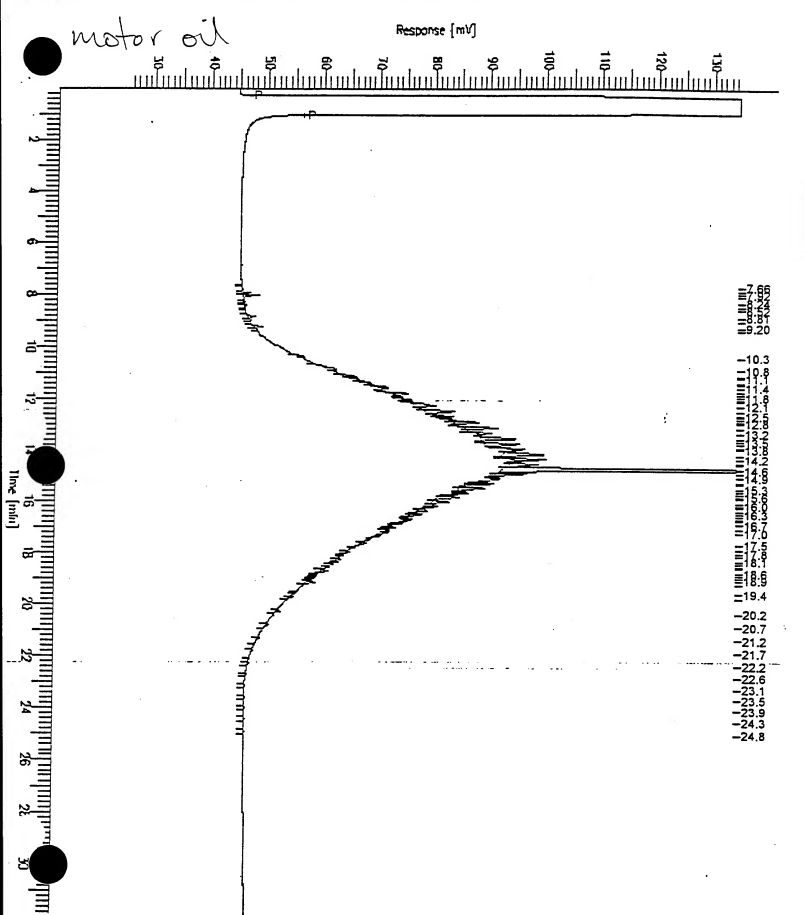
Time of Injection: 3/26/96 02:17 AM

Low Point : 25.07 mV

Plot Scale: 109.4 mV

High Point: 134.45 mV

Page 1 of 1



GC15 Channel A TEH

Sample Name : CCV, 96WS2069, JP5 FileName

: C:\GC15\CHB\084B023.RAW

Method : BTEH.MTH

Start Time : 0.01 min Scale Factor: 0.0

End Time : 19.80 min

Plot Offset: 42 mV

Sample #: 250MG/L

Date: 3/26/96 12:54 PM

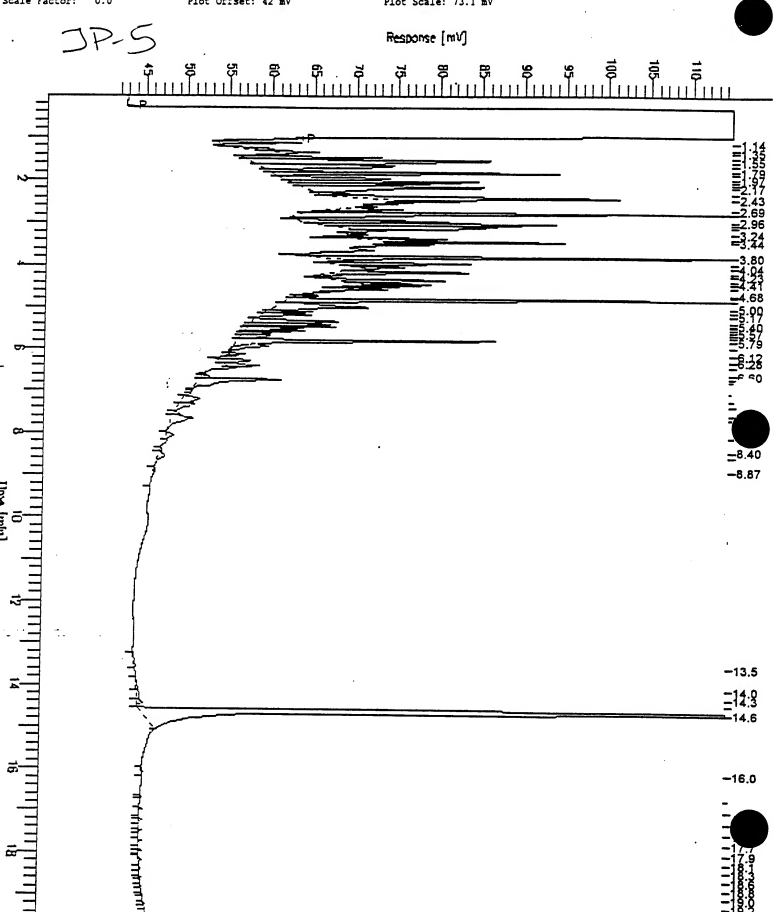
Time of Injection: 3/25/96 01:29 PM

Low Point: 41.54 mV Plot Scale: 73.1 mV

High Point : 114.59 mV

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TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124895-001 BE-IDW-01	26672	03/20/96	03/28/96	03/28/96	
124895-008 TRIP BLANK	26672	03/20/96	03/27/96	03/27/96	

Analyte Diln Fac:	Units	124895-001 1	124895-008 1	
Gasoline	ug/L	<50	<50	
Surrogate				
Trifluorotoluene	%REC	98	96	
Bromobenzene	%REC	101	94	



TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124895-007	COMP IDW-(2,3,4,5,6)	26608	03/20/96	03/25/96	03/25/96	17%

Analyte Diln Fac:	Units	124895-007 1	
Gasoline	mg/Kg	<1.2	
Surrogate			
Trifluorotoluene	%REC	98	
Bromobenzene	%REC	82	



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

client: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Method: EPA 5030

Location: Beale AFB

METHOD BLANK

 Matrix:
 Soil
 Prep Date:
 03/25/96

 Batch#:
 26608
 Analysis Date:
 03/25/96

Units: mg/Kg
Diln Fac: 1

MB Lab ID: QC17798

Analyte	Result	
Gasoline	<1.0	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene Bromobenzene	95 77	52-127 47-112



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: EPA 5030

METHOD BLANK

Matrix: Water Batch#: 26672 Units: ug/L Diln Fac: 1

Prep Date: 03/27/96 Analysis Date: 03/27/96

MB Lab ID: QC18055

Analyte	Result	
Gasoline	<50	•
Surrogate	%Rec	Recovery Limits
Trifluorotoluene Bromobenzene	95 87	69-120 70-122



BATCH QC REPORT

Page 1 of 1

Barria meraya a

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method:

EPA 5030

Location: Beale AFB

Soil

LABORATORY CONTROL SAMPLE

Prep Date:

03/25/96

26608 Batch#:

Matrix:

Analysis Date:

03/25/96

Units: mg/Kg Diln Fac: 1

LCS Lab ID: QC17799

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	9.6	10	96	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	112 92	52-127 47-112		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

 Matrix:
 Water
 Prep Date:
 03/27/96

 Batch#:
 26672
 Analysis Date:
 03/27/96

Units: ug/L Diln Fac: 1

LCS Lab ID: QC18056

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	1843	2006	92	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	103	69-120		
Bromobenzene	101	70-122		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits

BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Method: EPA 5030

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

 Field ID:
 ZZZZZZ
 Sample Date:
 03/12/96

 Lab ID:
 124797-006
 Received Date:
 03/13/96

 Matrix:
 Water
 Prep Date:
 03/27/96

 Batch#:
 26672
 Analysis Date:
 03/27/96

Units: ug/L Diln Fac: 1

MS Lab ID: QC18058

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Gasoline	2000	<50.00	2047	102	75-125
Surrogate	%Rec	Limits			
Trifluorotoluene	104	69-120			
Bromobenzene	110	70-122			

SD Lab ID: QC18059

nalyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Gasoline	2000	2135	107	75-125	4	<20
Surrogate	%Rec	Limit	s			-
Trifluorotoluene Bromobenzene	104 114	69-12 70-12	_	, "		

[#] Column to be used to flag recovery and RPD values with an asterisk

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits RPD: 0 out of 1 outside limits



Page 1 of

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020 Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124895-001 BE-IDW-01	26672	03/20/96	03/28/96	03/28/96	
124895-008 TRIP BLANK	26672	03/20/96	03/27/96	03/27/96	

Analyte Diln Fac:	Units	124895-001 1	124895-008 1	
Benzene	ug/L	<0.5	<0.5	
Toluene	ug/L	<0.5	<0.5	
Ethylbenzene	ug/L	<0.5	<0.5	
m,p-Xylenes	ug/L	<0.5	<0.5	
o-Xylene	ug/L	<0.5	<0.5	
1,2,3-Trimethylbenzene	∍ ug/L	<0.5	<0.5	
1,2,4-Trimethylbenzene	e ug/L	<0.5	<0.5	
1,3,5-Trimethylbenzene	e ug/L	<0.5	<0.5	
Surrogate				
Trifluorotoluene	%REC	98	95	
Bromobenzene	%REC	94	87	



BTXE

Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB

Prep Method:

Analysis Method: EPA 8020

EPA 5030

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124895-00	7 COMP IDW-(2,3,4,5,6)	26608	03/20/96	03/25/96	03/25/96	17%

Analyte Diln Fac:	Units	124895-007 1	
Benzene	ug/Kg	<6	
Toluene	ug/Kg	<6	
Ethylbenzene	ug/Kg	<6	
m,p-Xylenes	ug/Kg	<6	
o-Xylene	ug/Kg	<6	
1,2,3-Trimethylbenzer		<6	
1,2,4-Trimethylbenzer		<6	
1,3,5-Trimethylbenzer	e ug/Kg	<6	
urrogate			
Trifluorotoluene	%REC	100	
Bromobenzene	%REC	97	



BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc. Analysis Method: EPA 8020

Project#: 722450.14020 Location: Beale AFB

Prep Method: EPA 5030

METHOD BLANK

Matrix: Soil Batch#: 26608 Prep Date: Analysis Date:

03/25/96 03/25/96

Units: ug/Kg

Diln Fac: 1

MB Lab ID: QC17798

Analyte	Result	
Benzene	<5.0	
Toluene	<5.0	
Ethylbenzene	<5.0	
m, p-Xylenes	<5.0	
o-Xylene	<5.0	
1,2,3-Trimethylbenzene	<5.0	
1,2,4-Trimethylbenzene	<5.0	
1,3,5-Trimethylbenzene	<5.0	

Surrogate	%Rec	Recovery Limits
Trifluorotoluene .	99	43-114
Bromobenzene	94	45-140



BATCH QC REPORT

Page 1 of 1

		BTXE	
	Parsons Engineering Science, Inc. 722450.14020 Beale AFB	Analysis Method: Prep Method:	EPA 8020 EPA 5030
3.0	METI	HOD BLANK	
Matrix: Batch#: Units: Diln Fac:	Water 26672 ug/L	Prep Date: Analysis Date:	03/27/96 03/27/96

MB Lab ID: QC18055

Analyte	Result	
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	•
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene .	94	58-130
romobenzene	81	62-131



BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Soil Batch#: 26608 Units: ug/Kg Prep Date: 03/25/96 Analysis Date: 03/25/96

Diln Fac: 1

LCS Lab ID: QC17800

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	97	100	97	80-120
Toluene	101	100	101	80-120
Ethylbenzene	99	100	99	80-120
m,p-Xylenes	200	200	100	80-120
o-Xylene	103	100	103	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	100	43-114		
Bromobenzene	95	45-140		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 3 out of 8 outside limits



BATCH QC REPORT

Page 1 of 1

BTXE

lient: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

Project#: 722450.14020

Prep Method: EPA 5030

Aura Cara

Location: Beale AFB

LABORATORY CONTROL SAMPLE

03/27/96

Matrix: Water Batch#: 26672

Prep Date: Analysis Date: 03/27/96

Units: ug/L Diln Fac: 1

LCS Lab ID: QC18057

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	16.8	20	84	80-120
Toluene	17.7	20	89	80-120
Ethylbenzene	16.2	20	81	80-120
m,p-Xylenes	36.8	40	92	80-120
o-Xylene	17.5	20	88	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	96	58-130		
Bromobenzene	81	62-131		

Column to be used to flag recovery and RPD values with an asterisk lues outside of QC limits

ke Recovery: 0 out of 8 outside limits

124/895

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NIVIO	Curils & Tompkins, L.td. Analytical Laboratories, Since 1878 2323 Filth Street	(510) 486-0900 Phone (510) 486-0532 Fax Sn	0.14020	5, C	Tolect F.O.: 722450.14020 To		Sample ID. Date of the State of	-10w-01 12866000 -10w-02 0615	8E-102-25 K	10 P (no) Jac E			Sig

LOGIN CHANGE FOR	Date/Time: 3 21 9to Initials:	Matrix Add/Cancel Analysis	
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MPKINS, LTE	reguest Client Reques	Previous Lab ID	
CURTIS & TOI	Reason for change:	Current Lab ID	

Duedate											
Analysis	TWAR THURS			_		-		 	 		
Add/Cancel	Mdd	_									
Matrix	H,D										
Client ID	TRIP BLANK										
Previous Lab ID											
Current Lab ID	8-568/61										

Beule

COOLER RECEIPT CHECKLIST

Curtis & Tompkins, Ltc

	3 /20 #: 12485 Date Received: Number of Coolers:	1	
Login#			
Client:	Project: <u>Beull</u>		
A.	Preliminary Examination Phase	A	
_	Date Opened: 3/w By (print): Julia (sign)		_
1.	Did cooler come with a shipping slip (airbill, etc.)?	YES	NO
_	If YES, enter carrier name and airbill number:		
2.	Were custody seals on outside of cooler?	YES	MO
	How many and where? Seal date: Seal name:		
3.	Were custody seals unbroken and intact at the date and time of arrival?		•
4.	Were custody papers dry and intact when received?		NO
5.	Were custody papers filled out properly (ink, signed, etc.)?		NO
6.	Did you sign the custody papers in the appropriate place?		NO
7.	Was project identifiable from custody papers?		NO
	If YES, enter project name at the top of this form.		
8.	If required, was sufficient ice used? Type of ice: Temperature:	XE8	NO
	Type of ice: Cuhe / Shee Temperature: 4.75°	-	
	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \		
B.	Login Phase	$\Lambda_{1}\Lambda_{2}^{2}$	\cap i.
	Date Logged In: 3/20 By (print): 4William (sign)	<u> Jule</u>	<u> </u>
	Describe type of packing in cooler:		
	Did all bottles arrive unbroken?	(YES)	NO
3.	Were labels in good condition and complete (ID, date, time, signature, etc.)?		МО
	Did bottle labels agree with custody papers?		DD Jan
	Were appropriate containers used for the tests indicated?		NO
	Were correct preservatives added to samples?		NO
	Was sufficient amount of sample sent for tests indicated?		
	Were bubbles absent in VOA samples? If NO, list sample Ids below		
	Was the client contacted concerning this sample delivery?	YES	NO
	If YES, give details below.		
,	Who was called? By whom? Date):	
	mal Comments: Trinkluk received not an (-o-c down 3/20/a0)		
Addition	nal Comments:		
<u>*</u>	Trinkluk received not on (-o-c dow sio ao		
·			
	The Manager and a second		
-iiename: 🤄	:\qc\forms\cooler.wpd Re	v. 1 4/95	



Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 94710, Phone (510) 486-0900

ANALYTICAL REPORT

on the second of

Prepared for:

Parsons Engineering Science, Inc. 1301 Marina Village Parkway Suite 200 Alameda, CA 94501

Date: 02-APR-96

Lab Job Number: 124848

Project ID: 722450.14020

Location: Beale AFB

Reviewed by:

Reviewed by:

This package may be reproduced only in its entirety.



Laboratory Number: 124848 Sample Date: 03/15/96 Client: Parsons Engineering Science Receipt Date: 03/15/96

Project#: 722450.14020 Location: Beale AFB

CASE NARRATIVE

This hardcopy data package contains sample results and batch QC for four water samples received from the above referenced project on March 15, 1996. All samples were received cold and intact.

TPH/Extractable: No analytical problems were encountered.

TPH/Purgeable: No analytical problems were encountered.

BTXE (EPA 8020): No analytical problems were encountered.



TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Location: Beale AFB

Prep Method: EPA 3520

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124848-001	BE-SW01-01	26513	03/15/96	03/19/96	03/22/96	
124848-002	BE-SW02-01	26513	03/15/96	03/19/96	03/22/96	
124848-003	BE-SW12-01	26513	03/15/96	03/19/96	03/22/96	
124848-004	BE-SW03-01	26513	03/15/96	03/19/96	03/22/96	

Analyte Diln Fac:	Units	124848-001 1	124848-002 1	124848-003 1	124848-004 1
JP5	ug/L	<50	<50	<50	<50
Diesel Range	ug/L	<50	140 Y	120 Y	<50
Motor Oil Range	ug/L	<300	<300	<300	<300
Surrogate		1			
xacosane	%REC	104	127	120	108

Y: Sample exhibits fuel pattern which does not resemble standard



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Location: Beale AFB

Prep Method:

EPA 3520

METHOD BLANK

Matrix: Water Batch#: 26513 ug/L

Prep Date: 03/19/96 Analysis Date: 03/21/96

Units: Diln Fac: 1

MB Lab ID: QC17466

Analyte	Result	
JP5	<50	
Diesel Range	<50	
Motor Oil Range	<300	
Surrogate	%Rec	Recovery Limits
Hexacosane	103	60-140

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled	: 3/15/96	Client Project ID.	:	722450.14020
Date Received	: 3/16/96	Lab Project Number		96-0805
Date Prepared	: 3/16/96	Method		EPA 300.0
Date Analyzed	: 3/16/96	Detection Limit		0.056 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Nitrate-N</u> mg/L	Dilution <u>Factor</u>
X20559	BE-SW01-01	Water	0.15	1
X20560	BE-SW02-01	Water	0.12	1
X20561	BE-SW03-01	Water	0.17	1

Method Blank

(3/16/96)

< 0.056

Quality Assurance *

	<u>\$</u>	Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20559	BE-SW01-01 Matrix Spike	10.0	0.65	10.0	94
X20559	BE-SW01-01 Matrix Spike Duj	10.0	0.65	9.9	92
MS/MSD	RPD				1.6

Quality assurance results reported as Nitrate (NO₃).

Analyst

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled	: 3/15/96	Client Project ID.	:	722450.14020
Date Received	: 3/16/96	Lab Project Number	:	96-0805
Date Prepared	: 3/16/96	Method	:	EPA 300.0
Date Analyzed	: 3/16/96	Detection Limit	:	0.25 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Sulfate</u> mg/L	Dilution <u>Factor</u>
X20559	BE-SW01-01	Water	1.6	1
X20560	BE-SW02-01	Water	9.9	1
X20561	BE-SW03-01	Water	2.1	1

Method Blank

(3/16/96)

< 0.25

Quality Assurance

	:	Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20559	BE-SW01-01 Matrix Spike	10.0	1.6	10.9	94
X20559	BE-SW01-01 Matrix Spike Du	p 10.0	1.6	11.2	97
MS/MSD F	RPD				3.1

Analyst Analyst

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

 Date Sampled
 : 3/15/96
 Client Project ID.
 : 722450.14020

 Date Received
 : 3/16/96
 Lab Project Number
 : 96-0805

 Date Prepared
 : 3/16/96
 Method
 : EPA 300.0

 Date Analyzed
 : 3/16/96
 Detection Limit
 : 0.25 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Chloride</u> mg/L	Dilution <u>Factor</u>
X20559	BE-SW01-01	Water	2.3	1
X20560	BE-SW02-01	Water	33	5
X20561	BE-SW03-01	Water	5.1	1

Method Blank

(3/16/96)

< 0.25

Quality Assurance

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20559	BE-SW01-01 Matrix Spike	10.0	2.3	12.3	100
X20559	BE-SW01-01 Matrix Spike Du	ıp 10.0	2.3	14.3	120
MS/MSD F	RPD				18

Analyst

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled	: 3/15/96	Client Project ID.	:	722450.14020
Date Received	: 3/16/96	Lab Project Number	:	96-0805
Date Prepared	: 3/16/96	Method	:	EPA 300.0
Date Analyzed	: 3/16/96	Detection Limit	:	0.076 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Nitrite-N</u> mg/L	Dilution <u>Factor</u>
X20559	BE-SW01-01	Water	<0.076	1
X20560	BE-SW02-01	Water	<0.076	1
X20561	BE-SW03-01	Water	<0.076	1

Method Blank

(3/16/96)

< 0.076

Quality Assurance *

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20559	BE-SW01-01 Matrix Spike	10.0	<0.25	10.1	101
X20559	BE-SW01-01 Matrix Spike Du	p 10.0	<0.25	9.6	96
MS/MSD	RPD				5.5

^{• =} Quality assurance results reported as Nitrite (NO₂).

Amalyst Hol

Evergreen Analytical Sample Receipt/Ch			_
Date & Time Rec'd: 3/16/96 0927 Shipped Vi	a: F00	X-070	010914
Client: PARSON ES		. 4551104	51 0,
Client Project ID(s):			
EAL Project #(s):96-0805 EAL Co	oler(s):	Y	N
Cooler# PIRSONS			
Ice packs Y N Y N Y N Y	N	Y Y	
Temperature °C			-
·	Y	И	N/A
1. Custody seal(s) present: Seals on cooler intact Seals on bottle intact	X		
2. Chain of Custody present:	X		
3. Samples Radioactive: (Comment on COC if >0.5mr/b)		<u></u>	
4. Containers broken or leaking: (Comment on COCIFY)		X	
5. Containers labeled:	X		
6. COC agrees w/ bottles received: (Comment on COC # 5)	V		
7. COC agrees w/ labels: (Comment on COC UN)	V		
8. Headspace in vials-waters only: (Comment on COCIFY)			
9. VCA samples preserved:			
10. pH measured on metals, cyanide or phenolics*: List discrepancies *Non-EAL provided containers only, water samples			<u>~</u>
11. Metal samples present:	-		
Total, Dissolved, TCLP			
D or PD to be filtered: T,TR,D,PD to be Preserved:			
12. Short holding times: Specify parameters NOZ NOZ -			
13. Multi-phase sample(s) present:			
14. COC signed w/ date/time:	<u>/</u>		
Comments:			
(Additional comments on back)			
Custodian Signature/Date:	3-16-	96	

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

Date Sampled **Date Received**

: 3/15/96 : 3/16/96

Client Project ID.

: 722450.14020

Lab Project Number: 96-0805

Date Prepared

: 3/27/96

Method

: EPA 310.1

Date Analyzed

: 3/27/96

Detection Limit

: 5.0 mg CaCO₃/L

Evergreen Sample #	Client <u>Sample ID.</u>	<u>Matrix</u>	Total <u>Alkalinity</u> (mg CaCO ₃ /L)	Dilution <u>Factor</u>
X20559	BE-SW01-01	Water	42.2	1
X20560	BE-SW02-01	Water	92.2	1
X20561	BE-SW03-01	Water	40.3	1
X20562	BE-SW12-01	Water	93.2	1

Method Blank

(3/27/96)

< 5.0

Quality Assurance

Reference	True Value (mgCaCO ₃ /L)	<u>Result</u> (mgCaCO ₃ /L)	% Recovery
ERA Alkalinity Lot # 0814-95-02	120	122	102

Evergree	n Analytical S	ample Log Sheet	Project # <u>96-0805</u>
Date(s)	Sampled: 03/	15/96 COC	Date Due: 04/01/96
		/96 0927 Hold	Rush STANDARD
Address: Contact: Phone #	1301 MARINA ALAMEDA, CA MICHAEL PHEN 510-769-0100 Invoicing/Bil		Cooler Return 5.00 E.A. Cooler # N/A Airbill # FEDEX 0701091495 Client P.O. 722450.14020
Lab ID #	Client ID#	Analysis	Mtx Btl Loc
X20559A	BE-SW01-01	ALKALINITY	W 125P C6
X20560A	BE-SW02-01	ALKALINITY	W 125P C6
<u>561A</u>	BE-SW03-01	ALKALINITY	W 125P C6
X20562A	BE-SW12-01	ALKALINITY	W 125P C6
<u>у 9в</u>	BE-SW01-01	$NO_2, NO_3, SO_4, C1$	
X20560B	BE-SW02-01	NO ₂ , NO ₃ , SO ₄ , Cl ⁻	W 125P C6
X20561B	BE-SW03-01	NO ₂ ,NO ₃ ,SO ₄ ,Cl	W 125P C6
ate GC/	SxRec		
Page 1 of	1 Page(s)		Custodian/Date: 19/1/3/19/6

CHAIN OF CUSTODY RECORD / ANALYTICAL SERVICES REQUEST

7. 7.0 40							Eve	rgre	en A	nal	ytic	Evergreen Analytical Inc.	ပ							п.	Page_Lof_	
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2	1-0/00	ZIP 1130	2 ×	2/10	76	FAX# 510 - 769-924	77	,	88 8	(800) 845-7400 FAX RESULTS	300) 845-7400 FAX RESULTS	; ; ≻	(Z		EAL	EAL. QUOTE #	EAL. QUOTE #TURNAROUND REQUIRED*	IIBED.		P.O.# 7	PO.# 722450 14629	720
Sampler Name: (signature)	Buy								(I	#	Coc	1			exb.	edited	expedited turnaround subject to additional fee	d subjec	t to addi	ional fee		
(print) Michael	Phelps		Σ	MATRIX	×	40				A	NAL	YSIS	, RE	ANALYSIS REQUESTED	STE					EA Do	EAL use only Do not write	
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CLIENT	DATE	stnoO to	iter-Drink (circle)	bilos / I	egbul2 \	VAOV Cle) YA 8260/6	9/03S8 A(8 sepions	sVPCBs 8	B Screen	EX 8020/6	· · · · ·	-2108 Ha	600	M bevios Solved M Sie & list	מיסמ	1,70)4 1,47,7			Custodian		1
IDENTIFICATION	SAMPLED	TIME 2	_	ios		OT Gib	-	Α.		D d	18	_	∃T.	(Cir	Cir	V	ファ		_	EAL 9	EAL Sample No.	
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BE-5W02-01	3/15/96	1220 2	X												>	×					} -	
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BE-5W12-061	3/15/46	1230 8	X												X	4	John				62 4	T
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DD:						\dashv													8	Container Size	ize /25-	
Instructions: BE-5W12	10	is Alkalinity	٥(ر	14.	.	on (4																
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13(1577) CMM hand 1500

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nquished by: (Signature)

Date/Time Received by: (Signa

Date/Time

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

 Date Sampled
 : 3/13/96
 Client Project ID.
 : 722450.14020

 Date Received
 : 3/14/96
 Lab Project Number
 : 96-0776

 Date Prepared
 : 3/15/96
 Method
 : EPA 300.0

 Date Analyzed
 : 3/15/96
 Detection Limit
 : 0.25 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Sulfate</u> mg/L	Dilution <u>Factor</u>
X20416	BE-MP10-01	Water	12.2	1
X20417	BE-MP12-01	Water	21.1	1
X20418	BE-MP13-01	Water	1.1	1

Method Blank

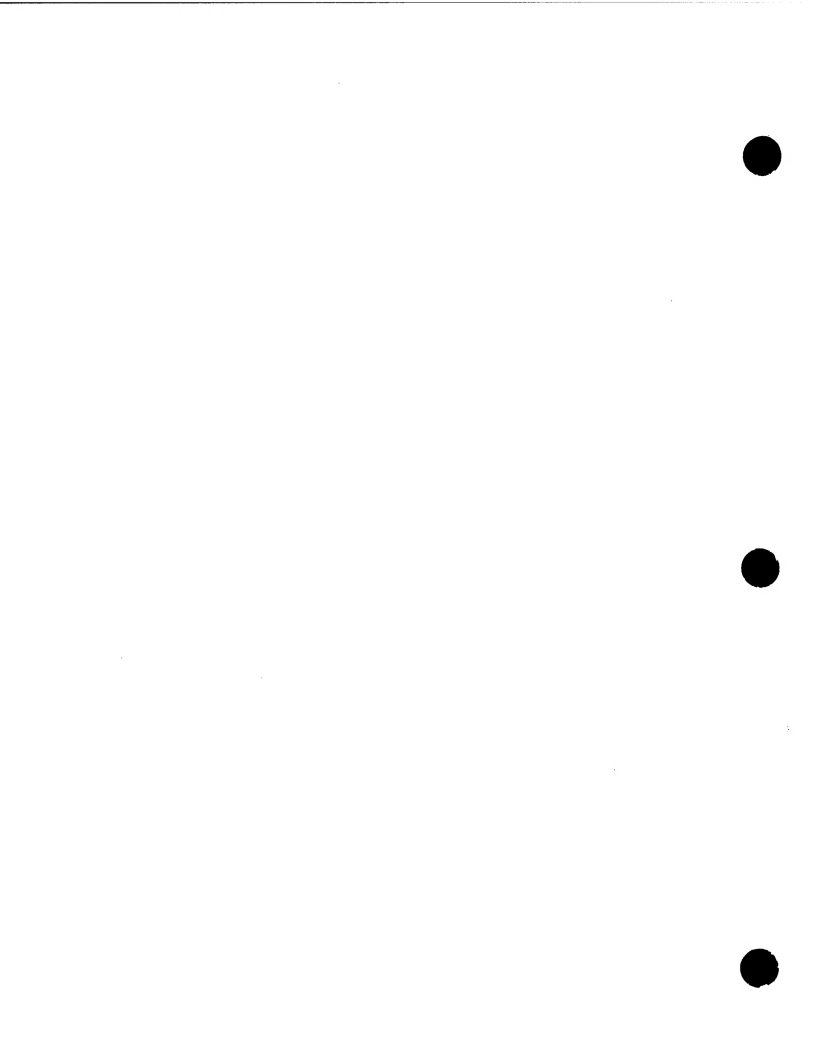
(3/15/96)

< 0.25

Quality Assurance

	<u>S</u> i	oike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20416	BE-MP10-01 Matrix Spike	10.0	12.2	21.2	91
X20416	BE-MP10-01 Matrix Spike Dup	10.0	12.2	22.0	98
MS/MSD	RPD				7.7

Analyst



4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

 Date Sampled
 : 3/13/96
 Client Project ID.
 : 722450.14020

 Date Received
 : 3/14/96
 Lab Project Number
 : 96-0776

 Date Prepared
 : 3/15/96
 Method
 : EPA 300.0

 Date Analyzed
 : 3/15/96
 Detection Limit
 : 0.076 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Nitrite-N</u> mg/L	Dilution <u>Factor</u>
X20416	BE-MP10-01	Water	<7.6 ⁽¹⁾	100
X20417	BE-MP12-01	Water	<0.076	1
X20418	BE-MP13-01	Water	<0.76 (1)	10

Method Blank

(3/15/96)

< 0.076

Quality Assurance *

	<u>\$</u>	pike Amount (mg/L)	Sample Result (2) (mg/L)	Spike Result (mg/L)	% Recovery
X20416	BE-MP10-01 Matrix Spike	10.0	<25	9.6	96
X20416	BE-MP10-01 Matrix Spike Dup	0 10.0	<25	9.6	96
MS/MSD F	RPD				0.62

^{* =} Quality assurance results reported as Nitrite (NO₂).

(2) = Sample result based on a 100X dilution factor due to matrix interferençe.

^{(1) =} Raised detection limit due to matrix interference.

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

: 3/13/96	Client Project ID.	:	722450.14020
: 3/14/96	Lab Project Number	:	96-0776
: 3/15/96	Method	:	EPA 300.0
: 3/15/96	Detection Limit	:	0.056 mg/L
	: 3/14/96 : 3/15/96	: 3/14/96	: 3/14/96

Evergreen Sample #	Client <u>Sample ID.</u>	<u>Matrix</u>	Nitrate-N mg/L	Dilution <u>Factor</u>
X20416	BE-MP10-01	Water	0.083	1
X20417	BE-MP12-01	Water	0.46	1
X20418	BE-MP13-01	Water	0.067	1

Method Blank

(3/05/96)

< 0.056

Quality Assurance *

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20416	BE-MP10-01 Matrix Spike	10.0	0.37	9.1	87
X20416	BE-MP10-01 Matrix Spike Du	p 10.0	0.37	9.6	92
MS/MSD F	RPD				5.9

Quality assurance results reported as Nitrate (NO₃).

Analyst

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

Date Sampled **Date Received**

: 3/13/96

Client Project ID.

: 722450.14020

: 3/14/96

Lab Project Number: 96-0776

Date Prepared Date Analyzed

: 3/20/96 : 3/20/96 Method

: Standard Method 4500-NH₃F

Detection Limit

: 0.80 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Ammonia-N	Dilution Factor
X20416	BE-MP10-01	Water	1.8 mg/L	1
X20417	BE-MP12-01	Water	<0.80 mg/L	1
X20418	BE-MP13-01	Water	<0.80 mg/L	1

.Method Blank

(3/20/96)

<0.80 mg/L

Quality Assurance

	Sı	oike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
<u>96-0741</u> X20229	Matrix Spike	10.0	<0.80	9.9	99
X20229	Matrix Spike Dup	10.0	<0.80	9.9	99
MS/MSD RF	סי				0.0

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled	: 3/13/96	Client Project ID.	:	722450.14020
Date Received	: 3/14/96	Lab Project Number	:	96-0776
Date Prepared	: 3/15/96	Method	:	EPA 300.0
Date Analyzed	: 3/15/96	Detection Limit	:	0.25 mg/L

Evergreen Sample #	Client <u>Sample ID.</u>	<u>Matrix</u>	<u>Chloride</u> mg/L	Dilution <u>Factor</u>
X20416	BE-MP10-01	Water	348	100
X20417	BE-MP12-01	Water	136	10
X20418	BE-MP13-01	Water	195	10

Method Blank

(3/15/96)

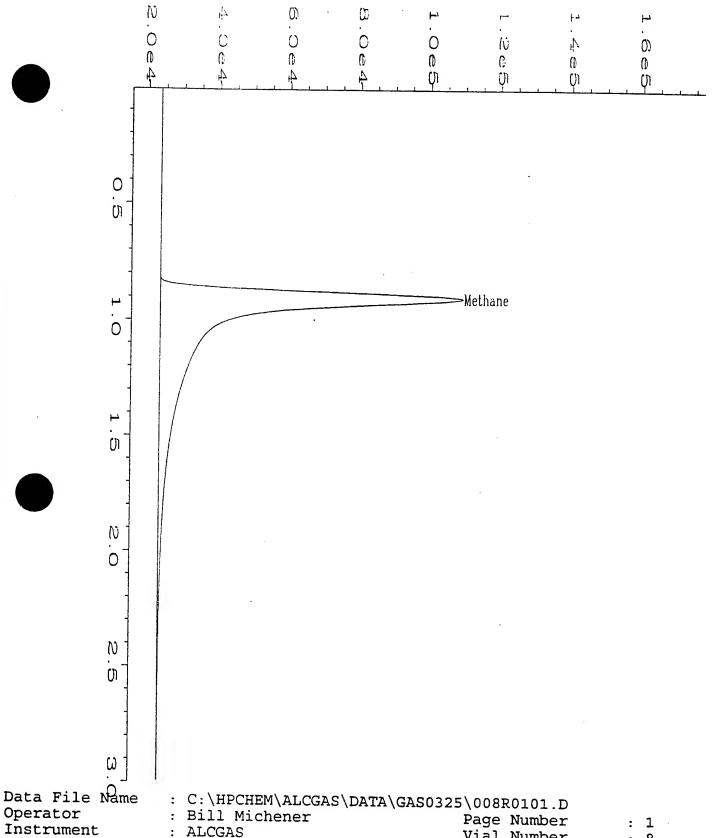
< 0.25

Quality Assurance

		Spike Amount (mg/L)	Sample Result * (mg/L)	Spike Result (mg/L)	% Recovery
X20416	BE-MP10-01 Matrix Spike	10.0	3.5	12.8	94
X20416	BE-MP10-01 Matrix Spike Du	p 10.0	3.5	12.6	92
MS/MSD F	RPD				2.3

^{* =} Sample result based on a 100X dilution factor.

///Analyst



Operator Instrument : ALCGAS Vial Number : 8 rple Name : LCS032596;Gas Injection Number: 1 Time Bar Code: Sequence Line red on : 25 Mar 96 09:57 AM Instrument Method: GAS.MTH Created on: 25 Mar 96 10:06 AM Analysis Method : GAS0325.MTH Last Recalib on : 25 MAR 96 09:44 AM Sample Amount : 0 Multiplier ISTD Amount Sample Info : Laboratory Control Sample Displaced 4ml of deionized water in 43ml vial with 1%

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

Date Sampled Date Received	: 3/13/96 : 3/14/96	Client Project ID.	
Date Prepared	: 3/14/96	Lab Project Number Method	: 96-0776 : EPA 310.1
Date Analyzed	: 3/18/96	Detection Limit	: $5.0 \text{ mg CaCO}_3/L$

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Total <u>Alkalinity</u> (mg CaCO ₃ /L)	Dilution <u>Factor</u>
X20416	BE-MP10-01	Water	690	1
X20417	BE-MP12-01	Water	213	1
X20418	BE-MP13-01	Water	575	1

Method Blank

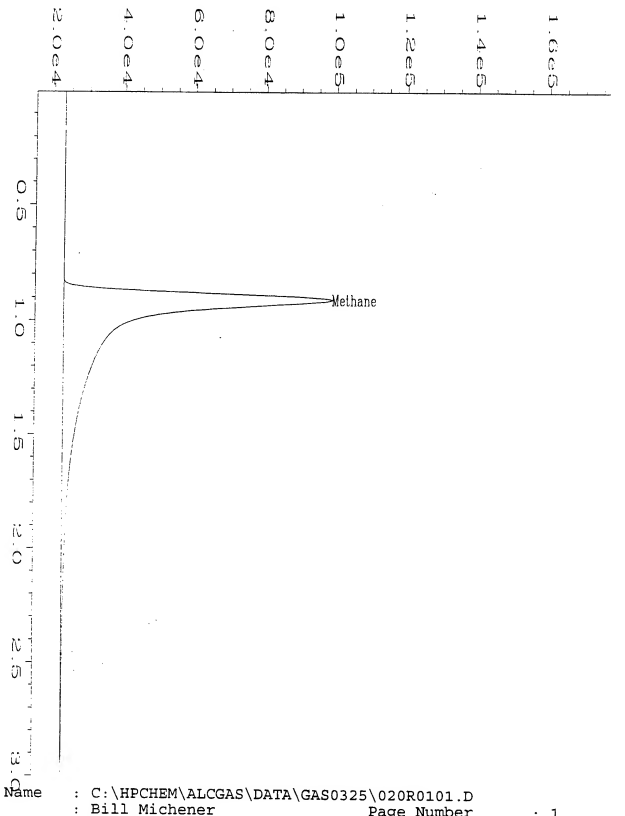
(3/18/96)

< 5.0

Quality Assurance

Reference	<u>True Value</u> (mgCaCO ₃ /L)	<u>Result</u> (mgCaCO ₃ /L)	% Recovery
ERA Alkalinity Lot #	120	116	97

Analyst Ope



Data File Name :	C:\HPCHEM\ALCGAS\DATA\GAS032	25\020R0101.D
Operator :	Bill Michener	Page Number : 1
Instrument :	ALCGAS	Vial Number : 20
	X20417MSD;1	Injection Number : 1
Run Time Bar Code:	, <u> </u>	Sequence Line : 1
	25 Mar 96 01:37 PM	Instrument Method: GAS.MTH
Actort Created on:	25 Mar 96 01:42 PM	Analysis Mothod GASONIA
La Recalib on :	25 MAR 96 09:44 AM	Analysis Method : GAS0325.MTH Sample Amount : 0
	1	
		ISTD Amount :
pambre into :	96-0776;BE-MP12-01;Displaced	4ml with 1% methane
	gas($\#1/23$), shook for 5 min.	and injected 50ul to equal a
	theoretical spike of 500mg	The sample is injected at a

Evergreen Analytical, Inc. 4036 Youngfield, Wheat Ridge, CO 80033 (303) 425-6021

RSK-175 Gas Method Methane LCS Report Form

LCS No.

: LCS032596

EPA Method No.

: RSKSOP-175

Date Prepared

: 3/25/96

Matrix

: Water

Date Analyzed

: 3/25/96

Method Blank

: GB032596

E.A. LCS Source No.

: 1723

Lab File No.

: GAS0325008

	Spike	Method Blank	LCS		QC
Compound	Added	Concentration	Concentration	LCS	Limits
	(ug)	(ug)	(ug)	%REC	%REC
Methane Gas	500	0	407	81	67-85

Spike Recovery:

0

0 out of (1) outside limits.

Note: The LCS was made by taking the sample and displacing 4ml of headspace with a 1% methane gas and shaking the VOA for 5 minutes. Then injecting 50 ul from the headspace into the GC resulting in a theoretical concentration of 500 ug.

NOTES:

* = Values outside of QC limits.

NA = Not analyzed/not available.

Analyst

Approved

LCS0325.XLS; 3/25/96

Evergreen Analytical, Inc. 4036 Youngfield, Wheat Ridge, CO 80033 (303) 425-6021

RSK-175 Gas Method Methane, Ethane, Ethene Gas Matrix Spike / Matrix Spike Duplicate Report

Client Sample No.

: BE-MP12-01

Client Project No.

: 722450.14020

Lab Sample No.

: X20417

Lab Project No.

: 96-0776

Date Sampled

: 3/13/96

EPA Method No.

: RSKSOP-175

Date Received

: 3/14/96

Matrix

: Water

Date Prepared

: 3/25/96

Method Blank

: GB032596

Date Analyzed

: 3/25/96

Lab File No's.

: GAS0325019,020

E.A. MS/MSD Spike Source No.

: 1723

	Spike	Sample	MS		QC
Compound	Added	Concentration	Concentration	MS	Limits
	(ug)	(ug)	(ug)	%REC	%REC
Methane Gas	500	0	379	76	40-89

	Spike	MSD				C
Compound	Added	Concentration	MSD	RPD	Lin	nits
	(ug)	(ug)	%REC		RPD	%REC
Methane Gas	500	382	76	0.8	0-24.4	40-89

out of (1) outside limits. out of (2) outside limits.

Spike Recovery:

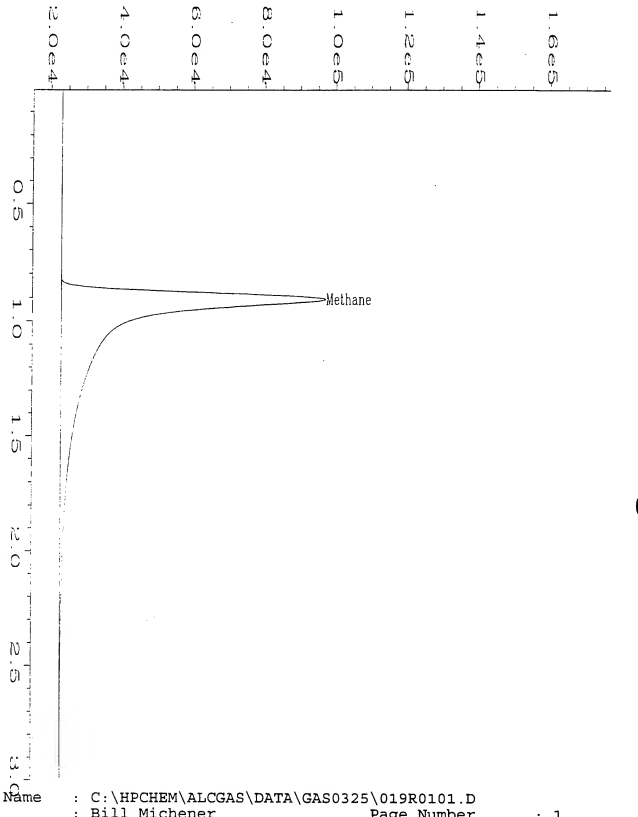
NOTES:

* = Values outside of QC limits.

NA = Not analyzed/not available

Note: The Spike was made by taking the sample and displacing 4ml of headspace with a 1% methane gas and shaking the VOA for 5 minutes. Then injecting 50 ul from the headspace into the GC resulting in a theoretical concentration of 500 ug.

MS0776.XLS; 3/25/96



Data File Name Operator : Bill Michener Page Number : 1 Instrument : ALCGAS Vial Number : 19 Sample Name : X20417MS;1 Injection Number: 1 Run Time Bar Code: Sequence Line : 1 Acquired on : 25 Mar 96 01:33 PM Instrument Method: GAS.MTH Report Created on: 25 Mar 96 01:42 PM Analysis Method : GAS0325.N Last Recalib on : 25 MAR 96 09:44 AM Sample Amount Multiplier ISTD Amount Sample Info : 96-0776; BE-MP12-01; Displaced 4ml with 1% methane gas(#1723), shook for 5 min. and injected 50ul to equal a

theoretical spike of 500mg. The sample is injected at a

Methane Report Form

t Sample Number	: BE-MP13-01	Client Project No.	: 722450.14020
Sample Number	: X20418	Lab Project No.	: 96-0776
Date Sampled	: 3/13/96	Dilution Factor	: 50.00
Date Received	: 3/14/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/25/96	Matrix	: Water
Date Analyzed	: 3/25/96	Lab File No.	: GAS0325017

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	1.6	0.1

mperature	: _	75 F	Saturation	Meth	0.38539415
nount Injected	: _	0.01 ml	Concentration		
Volume of Sample	: _	43 ml	Concentration	Meth	1.207475299
space created	; _	4 ml	in Head Space		
Methane Area	: _	179.243 ug			

16 g

QUALIFIERS:

E = Extrapolated value.

Atomic weight(Methane)

U = Compound analyzed for, but not detected.

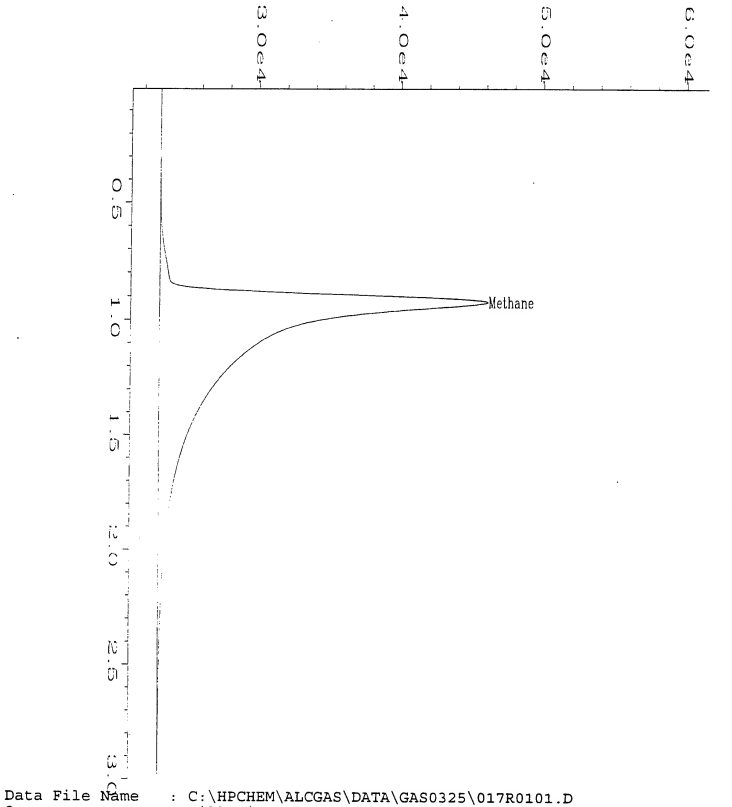
B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

At 28 pproved

AF0776.XLS



Operator			Page Number :	1
Instrument	:	ALCGAS	Vial Number :	
Sample Name	:	X20418;50	Injection Number :	
Run Time Bar Code			Sequence Line :	
Acquired on	:	25 Mar 96 01:09 PM	Instrument Method:	
Report Created on	1:	25 Mar 96 01:42 PM	Analysis Method :	
Last Recalib on	:	25 MAR 96 09:44 AM	Sample Amount :	
Multiplier	:	1	ISTD Amount :	
Sample Info	:	96-0776;BE-MP13-01;Water	•	

Methane Report Form

1020
75
16

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

nperature	:	73.8 F	Saturation	Meth	0.000108839
nount Injected	:	0.5 ml	Concentration		
Volume of Sample	:	43 ml	Concentration	Meth	0.00034177
space created	:	4 ml	in Head Space		
Methane Area	:	2.531 ug			

Atomic weight(Methane) : _____ g

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Approved

AF0776.XLS

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2.3e4
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                                              2.5c4
                                                              2.6e4
                                                                              2.7e4
                                                                                              2.8e4
                                                                                                                                               3.1e4
                                                                                                               2.9e4
                                                                                                                               3.0e4
                                                                                                                                                                3.2e4
 0.5
                           >Methane
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Data File Name :	C:\HPCHEM\ALCGAS\DATA\GAS0325	5\016R0101.D	
Operator :	Bill Michener	Page Number :	1
Instrument :		Vial Number :	
Sample Name :	X20417;1	Injection Number :	
Run Time Bar Code:		Sequence Line :	
Acquired on :	25 Mar 96 12:59 PM	Instrument Method:	
Report Created on:		Analysis Method :	
Last Recalib on :		Sample Amount :	
	1	ISTD Amount :	,

Sample Info : 96-0776;BE-MP12-01;Water

Methane Report Form

t Sample Number	: BE-MP10-01	Client Project No.	: 722450.14020
Sample Number	: X20416	Lab Project No.	: 96-0776
Date Sampled	: 3/13/96	Dilution Factor	: 20.00
Date Received	: 3/14/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/25/96	Matrix	: Water
Date Analyzed	: 3/25/96	Lab File No.	: GAS0325015

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	0.85	0.04

mperature	:	72.6 F	Saturation	Meth	0.204572839
.nount Injected	:	0.025 ml	Concentration		
Volume of Sample	:	43 ml	Concentration	Meth	0.643836992
space created	:	4 ml	in Head Space		
Memane Area	:	237.862 ug			

QUALIFIERS:

E = Extrapolated value.

Atomic weight(Methane)

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

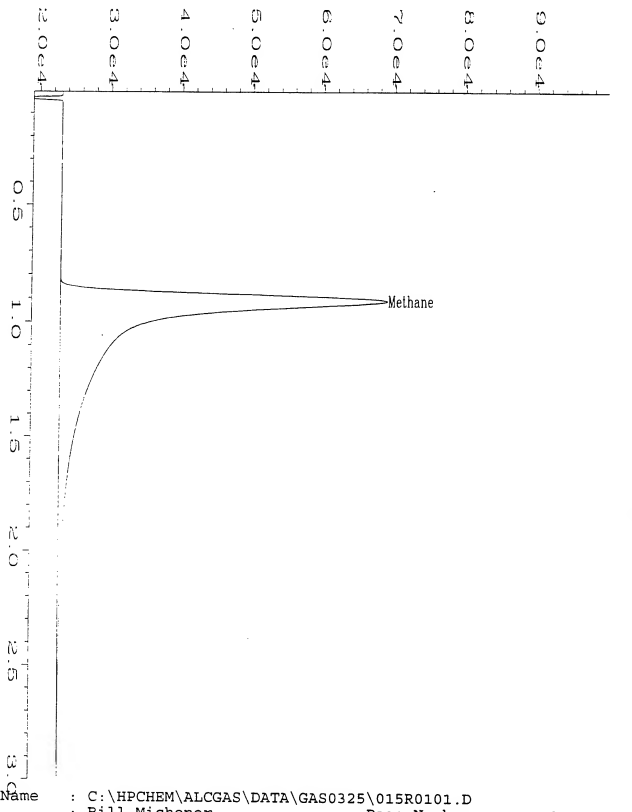
RL = Reporting Limit.

NA = Not Available/Not Applicable.

Millioner Analyst

Approved

AF0776.XLS



Data File Name	:	C:\HPCHEM\ALCGAS\DATA\GAS032	5\015R0101.D
Operator	:	Bill Michener	Page Number : 1
Instrument			Vial Number : 15
Sample Name	:	X20416;20	Injection Number : 1
Run Time Bar Code			Sequence Line : 1
Acquired on	:	25 Mar 96 12:46 PM	Instrument Method: GAS.MTH
Report Created on	:	25 Mar 96 01:42 PM	Analysis Method : GAS0325.M2
		25 MAR 96 09:44 AM	Sample Amount : 0
		1	ISTD Amount :
Sample info	:	96-0776;BE-MP10-01;Water	

Methane Report Form Method Blank Report

method Blank Number

: GB032596

Client Project No.

: 722450.14020

Date Extracted/Prepared

: 3/25/96

Lab Project No.

: 96-0776

Date Analyzed

: 3/25/96

Dilution Factor

: 1.00

Method

: RSKSOP-175

Matrix

: Water

Lab File No.

: GAS0325002

Sample

Compound Name

Cas Number

Concentration mg/L

RL mg/L

Methane

74-82-8

U

0.002

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Approved

AF0776.XLS

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Data File Name
                         : C:\HPCHEM\ALCGAS\DATA\GAS0325\002R0101.D
```

Operator : Bill Michener Page Number : 1 Instrument : ALCGAS Vial Number Sample Name : GB032596 Injection Number : 1 Run Time Bar Code: Sequence Line : 1 Acquired on : 25 Mar 96 08:51 AM Instrument Method: GAS.MTH Report Created on: 25 Mar 96 09:45 AM Analysis Method : GAS0325.N Last Recalib on : 25 MAR 96 09:44 AM Sample Amount : 0 Multiplier : 1 ISTD Amount Sample Info : Gas Method Blank

Displaced 4ml of deionized water in 43ml vial with Helium, shook for 5 min and injected 500ml

Evergreen Apalypical sample Receipt/Check-in R	,80072
Date & Time Rec'd: 3 14 9t 0925 Shipped Via: Fed	1154092520
Client: Yarsons ES	
Client Project ID(s): 722450, 14020	
EAL Project #(s):96-0776	i (i) n
Cooler# <u>735</u>	
Ise packs 💯 N Y N Y N Y N	ā ā
Temperature of 5	
. 7	N N/A
1. Custody seal(s) present: Seals on cocler intact Seals on bottle intact	
2. Chain of Custody present:	
3. Samples Radicactive: (Comment on COCT > (Limits)	X_
4. Containers broken or leaking: Comment on COC 479	×
5. Containers labeled:	
6. COC agrees w/ bottles received: Comment in COCIN ROC'd why & bettles for BE-MP12-01- No alkalinity 7. COC agrees w/ labels: Comment in COCIN	bottle rec'd - 1949.79
8. Headspace in vials-waters only: Commence cooks	<u> </u>
9. VOA samples preserved:	
10. pH measured on metals, cyanide or phenolics*: List discrepancies *Non-EAL provided containers only, water samples only.	
11. Metal samples present: Total, Dissolved, TCLP D or PD to be filtered: T,TR,D,PD to be Preserved:	
12. Short holding times: Specify parameters Now NO3	
13. Multi-phase sample(s) present:	K
14. COC signed w/ date/time:	
Comments:	
(Additional comments on back)	
Custodian Signature/Date: 12/19/96	

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Total Organic Carbon

Date Sampled : 3/13/96 Date Received : 3/14/96 Date Prepared : 3/15/96 Date Analyzed : 3/15/96	Lab Project Number Method	:	722450.14020 96-0776 EPA 415.1 1.0 mg C/L
--	------------------------------	---	--

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	TOC mg C/L	Dilution Factor
X20416	BE-MP10-01	Water	65.7	1
X20417	BE-MP12-01	Water	5.2	1
X20418	BE-MP13-01	Water	36.8	1

Method Blank

(3/15/96)

<1.0

Quality Assurance

	\$	Spike Amount (mgC/L)	Sample Result (mgC/L)	Spike Result (mgC/L)	% Recovery
X20416	BE-MP10-01 Matrix Spike	10.0	65.7	76.8	112
X20416	BE-MP10-01 Matrix Spike Dup	10.0	65.7	77.6	119
MS/MSD	RPD				6.4

/// Hal-

Evergree	n Analytical Sa	ample Log Sheet	Project	# <u>96-0776</u>
Date(s)	Sampled: 03/1	.3/96 COC	Date Due:	03/28/96
		<u>722450.14020</u>	Holding Time(s):	03/15-NO ₂ , NO ₃ , 03/27-ALKALINITY Rush STANDARD
Client:	PARSONS ENGIN	EERING SCIENCE, INC.	Cooler Return	n N/A
Address:	1301 MARINA	VILLAGE PKWY #200	E.A. Cooler;	# <u>735</u>
	ALAMEDA, CA	94501	Airbill # FEI	DEX 1154092520
Contact:	MICHAEL PHEL	PS	Client P.O.	1
Phone #	510-769-0100	Fax #510-769-9244		
Special 1	Invoicing/Bil	ling		
phecial 1	rns cruc crons_			
Lab ID #	Client ID#	Analysis	Mtx	Btl Loc
X20416A	BE-MP10-01	TOC	W	125A CR3
X20417A	BE-MP12-01	TOC	W	125A CR3
٧2 <u>0418A</u>	BE-MP13-01		W	125A CR3
	BE-MP10-01	NO ₂ , NO ₃ , SO ₄ , Cl	_	125P CR3
X2 7B	BE-MP12-01			125P CR3
X218B	BE-MP13-01			125P CR3
X20416C	BE-MP10-01	AMMONIA		500P CR3
X20417C	BE-MP12-01	AMMONIA	W	500P CR3
X20418C	BE-MP13-01	AMMONIA	W	500P CR3
X20416D	BE-MP10-01	ALKALINITY	W	500P CR3
X20418D	BE-MP13-01	ALKALINITY	W	500P CR3
<u> X20416E-G</u>	BE-MP10-01	METHANE	W	40V CR3
X20417E/F	BE-MP12-01	METHANE	W	40V CR3
X20418E/F	BE-MP13-01	METHANE	W	40V CR3
·				
		•		
R=Sample	to be returne	ed.		
te GC/	MS GC	X Metals Wes	t Chem X HPLC	SxPrep
	SxRec	C QA/QC C Accto	g <u>C</u> File <u>(</u>	Orig C
Page 1 of	1 Page(s)		Custodia	1/20/2/2
			Castoula	Am 2/21/91
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CHAIN OF CUSTODY RECORD / ANALYTICAL SERVICES REQUEST

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COMPANY PHESONS	s Ex	١.			•				4	036 Yo	4036 Youngfield St	ı Jor	}			CLIEN	S S S	TACT (print	Z	CLIENT CONTACT (print) LICETACA	PATAR	,
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Sampler Name: HILACH ECH F	ECH LE	. 1				•											(30			ō	€		
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nature) | 3 Date/Time Received by: (Signature)

·linquished by: (Signature)

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled : 3/12/96
Date Received : 3/13/96
Date Prepared : 3/14/96
Date Analyzed : 3/14/96

Client Project ID. : 722450.14020

Lab Project Number : 96-0741

Method : EPA 300.0

Detection Limit : 0.056 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Nitrate-N</u> mg/L	Dilution <u>Factor</u>
X20226	BE-MP22-01	Water	1.8	1
X20227	BE-MP42-01	Water	1.9	1
X20228	BE-MP11-01	Water	0.81	1
X20229	BE-MP44-01	Water	0.80	1

Method Blank

(3/14/96)

Water

< 0.056

Quality Assurance *

	<u>Sr</u>	oike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20226	Matrix Spike	10.0	8.2	18.6	104
X20226	Matrix Spike Dup	10.0	8.2	18.0	98
MS/MSD	RPD .				5.7

 ⁼ Quality assurance results reported as Nitrate (NO₃).

Analyst For

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled Date Received	: 3/12/96 : 3/13/96	Lab Project Number	:	
Date Prepared Date Analyzed	: 3/14/96 : 3/14/96	Method Detection Limit	•	EPA 300.0 0.25 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Sulfate mg/L	Dilution <u>Factor</u>
X20226	BE-MP22-01	Water	23.5	1
X20227	BE-MP42-01	Water	23.5	1
X20228	BE-MP11-01	Water	7.0	1
X20229	BE-MP44-01	Water	7.0	1

Method Blank

(3/14/96)

Water

< 0.25

Quality Assurance

	<u>:</u>	Spike Amount (mg/L)	Sample Result * (mg/L)	Spike Result (mg/L)	% Recovery
X20226	Matrix Spike	10.0	2.5	11.7	93
X20226	Matrix Spike Du	p 10.0	2.5	11.7	93
MS/MSD	RPD				0.11

^{* =} Value based on a 10X dilution factor.

Analyst

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled : 3/12/96 **Date Received** : 3/13/96

Client Project ID. Lab Project Number: 96-0741

: 722450.14020

Date Prepared Date Analyzed

: 3/14/96 : 3/14/96 Method

: EPA 300.0

Detection Limit

: 0.25 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Chloride</u>	Dilution <u>Factor</u>
X20226	BE-MP22-01	Water	72.2 mg/L	10
X20227	BE-MP42-01	Water	70.2 mg/L	10
X20228	BE-MP11-01	Water	18.0 mg/L	1
X20229	BE-MP44-01	Water	16.8 mg/L	1

Method Blank

(3/14/96)

Water

<0.25 mg/L

Quality Assurance

	<u> </u>	Spike Amount (mg/L)	Sample Result * (mg/L)	Spike Result (mg/L)	% Recovery
X20226	Matrix Spike	10.0	7.2	16.6	93
X20226	Matrix Spike Du	p 10.0	7.2	16.6	93
MS/MSD F	RPD				0

⁼ Value based on a 10X dilution factor.

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled : 3	3/12/96	Client Project ID. :	722450.14020
Date Received : 3	3/13/96	Lab Project Number:	96-0741
Date Prepared : 3	3/14/96	Method:	EPA 300.0
Date Analyzed : 3	3/14/96	Detection Limit :	0.076 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Nitrite-N</u> mg/L	Dilution <u>Factor</u>
X20226	BE-MP22-01	Water	<0.076	1
X20227	BE-MP42-01	Water	<0.076	1
X20228	BE-MP11-01	Water	<0.076	1
X20229	BE-MP44-01	Water	<0.076	1
Method Blank	(3/14/96)	Water	< 0.076	

Quality Assurance *

	<u>S</u>	pike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20226	Matrix Spike	10.0	<0.25	9.5	95
X20226	Matrix Spike Dup	10.0	<0.25	8.9	89
MS/MSD	RPD				5.9

^{* =} Quality assurance results reported as Nitrite (NO₂).

Analyst Hall

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

Date Sampled : 3/12/96 **Date Received** : 3/13/96 Date Prepared : 3/18/96 Date Analyzed : 3/18/96

Client Project ID.

: 722450.1420

Lab Project Number: 96-0741 Method

: EPA 310.1

Detection Limit

: 5.0 mg CaCO₃/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Total <u>Alkalinity</u> (mg CaCO ₃ /L)	Dilution <u>Factor</u>
X20226	BE-MP22-01	Water	181	1
X20227	BE-MP42-01	Water	186	1
X20228	BE-MP11-01	Water	104	1
X20229	BE-MP44-01	Water	103	1

Method Blank

(3/18/96)

< 5.0

Quality Assurance

Reference	<u>True Value</u> (mgCaCO ₃ /L)	<u>Result</u> (mgCaCO ₃ /L)	% Recovery
ERA Alkalinity Lot # 0814-95-06	120	116	97

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

 Date Sampled
 : 3/12/96
 Client Project ID.
 : 722450.1420

 Date Received
 : 3/13/96
 Lab Project Number
 : 96-0741

Date Prepared : 3/20/96 Method : Standard Method 4500-NH₃F

Date Analyzed : 3/20/96 Detection Limit : 0.80 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Ammonia-N	Dilution <u>Factor</u>
X20226	BE-MP22-01	Water	<0.80 mg/L	1
X20227	BE-MP42-01	Water	<0.80 mg/L	1
X20228	BE-MP11-01	Water	<0.80 mg/L	1
X20229	BE-MP44-01	Water	<0.80 mg/L	1

Method Blank (3/20/96)

<0.80 mg/L

Quality Assurance

	<u>s</u>	pike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20229	BE-MP44-01 Matrix Spike	10.0	<0.80	9.9	99
X20229	BE-MP44-01 Matrix Spike Dup	10.0	<0.80	9.9	99

MS/MSD RPD

0.0

Analyst

Evergreen Analytical, Inc. 4036 Youngfield, Wheat Ridge, CO 80033 (303) 425-6021

RSK-175 Gas Method Methane LCS Report Form

LCS No.

: LCS032596

EPA Method No.

: RSKSOP-175

Date Prepared

: 3/25/96

Matrix

: Water

Date Analyzed

: 3/25/96

Method Blank

: GB032596

E.A. LCS Source No.

: 1723

Lab File No.

: GAS0325008

	Spike	Method Blank	LCS		QC
Compound	Added	Concentration	Concentration	LCS	Limits
	(ug)	(ug)	(ug)	%REC	%REC
Methane Gas	500	0	407	81	67-85

Spike Recovery: 0 out of (1) outside limits.

Note: The LCS was made by taking the sample and displacing 4ml of headspace with a 1% methane gas and shaking the VOA for 5 minutes. Then injecting 50 ul from the headspace into the GC resulting in a theoretical concentration of 500 ug.

NOTES:

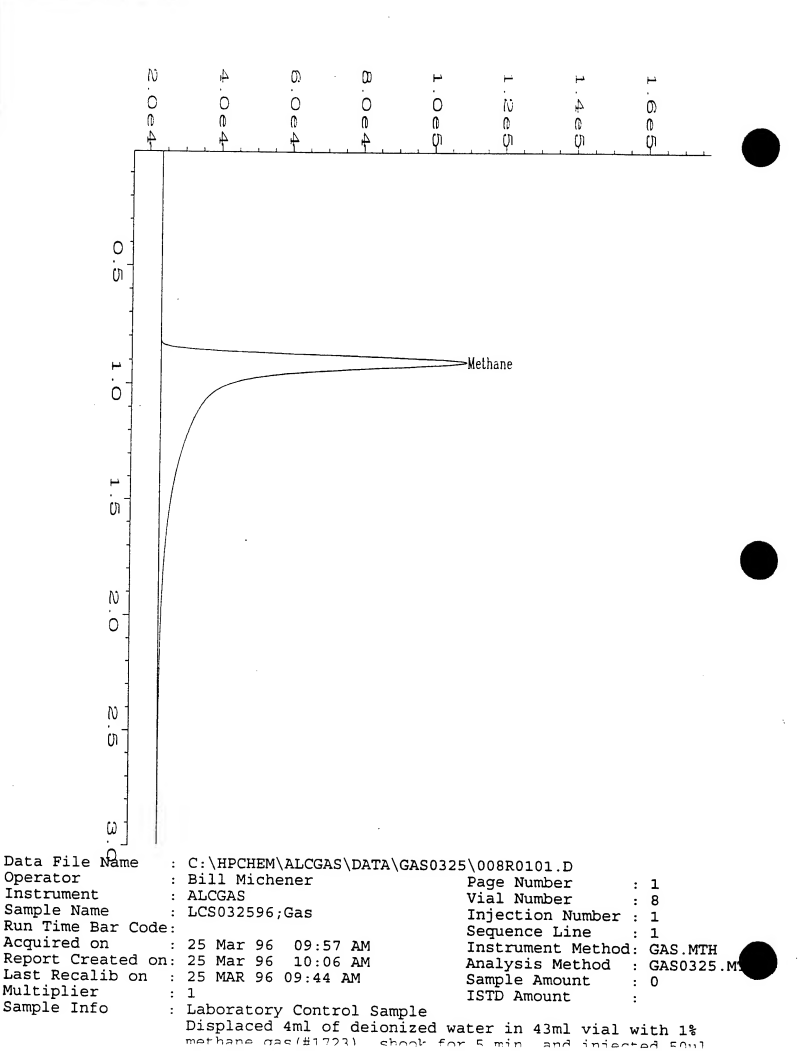
* = Values outside of QC limits.

NA = Not analyzed/not available.

Analyst Melsonor

Approved

LCS0325.XLS; 3/25/96



Methane Report Form

Sample Number ample Number Date Sampled Date Received Date Extracted/Prepared Date Analyzed	: BE-MP44-01 : X20229Dup : 3/12/96 : 3/13/96 : 3/25/96 : 3/25/96	Client Project No. Lab Project No. Dilution Factor Method Matrix Lab File No.	: 722450.14020 : 96-0741 : 1.00 : RSKSOP-175 : Water : GAS0325014
Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	0.164	0.002
ກperature ount Injected	: <u>72.1</u> : 0.5		Meth 0.03944152
Total Volume of Sample Pace created Memore Area	: 43	ml Concentration M ml in Head Space	Meth 0.124248158

<u>16</u> g

QUALIFIERS:

E = Extrapolated value.

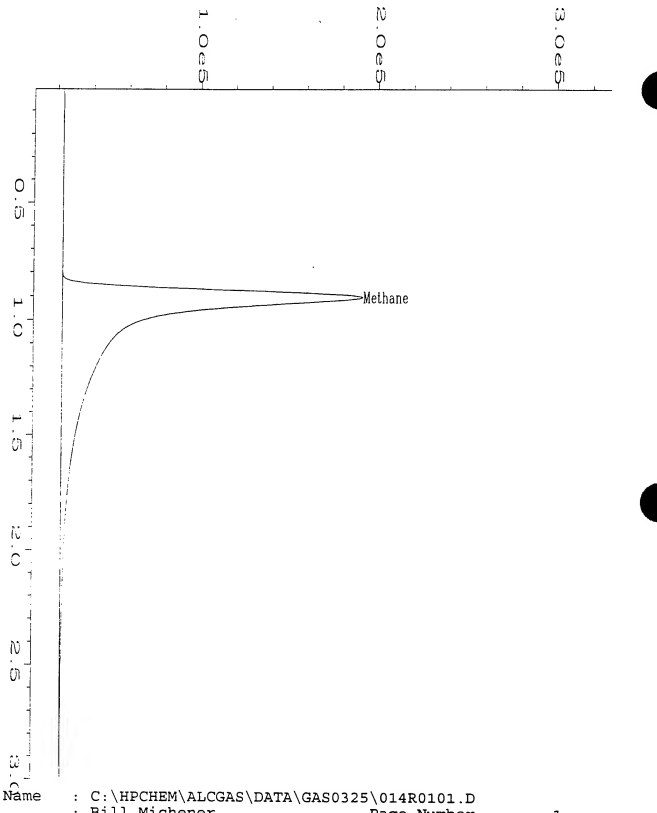
Atomic weight(Methane)

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.



Data File Name Operator : Bill Michener Page Number 1 Instrument : ALCGAS Vial Number 14 Sample Name : X20229Dup;1 Injection Number Run Time Bar Code: Sequence Line 1 Acquired on : 25 Mar 96 Instrument Method: GAS.MTH 12:29 PM Report Created on: 25 Mar 96 01:42 PM Analysis Method : GAS0325.M Last Recalib on : 25 MAR 96 09:44 AM Sample Amount : 0 Multiplier ISTD Amount

Sample Info : 96-0741;BE-MP44-01;Water

Methane Report Form

Sample Number	: BE-MP44-01	Client Project No.	: 722450.14020
Sample Number	: X20229	Lab Project No.	: 96-0741
Date Sampled	: 3/12/96	Dilution Factor	: 1.00
Date Received	: 3/13/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/25/96	Matrix	: Water
Date Analyzed	: 3/25/96	Lab File No.	: GAS0325013

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	0.167	0.002

mperature	:	72.1 F	Saturation	Meth _	0.040309739
Jount Injected	;	0.5 ml	Concentration		
Total Volume of Sample	:	43 ml	Concentration	Meth	0.12698321
pace created	:	4 ml	in Head Space		
Memane Area	:	937.383 ug			

16 g

QUALIFIERS:

E = Extrapolated value.

Atomic weight(Methane)

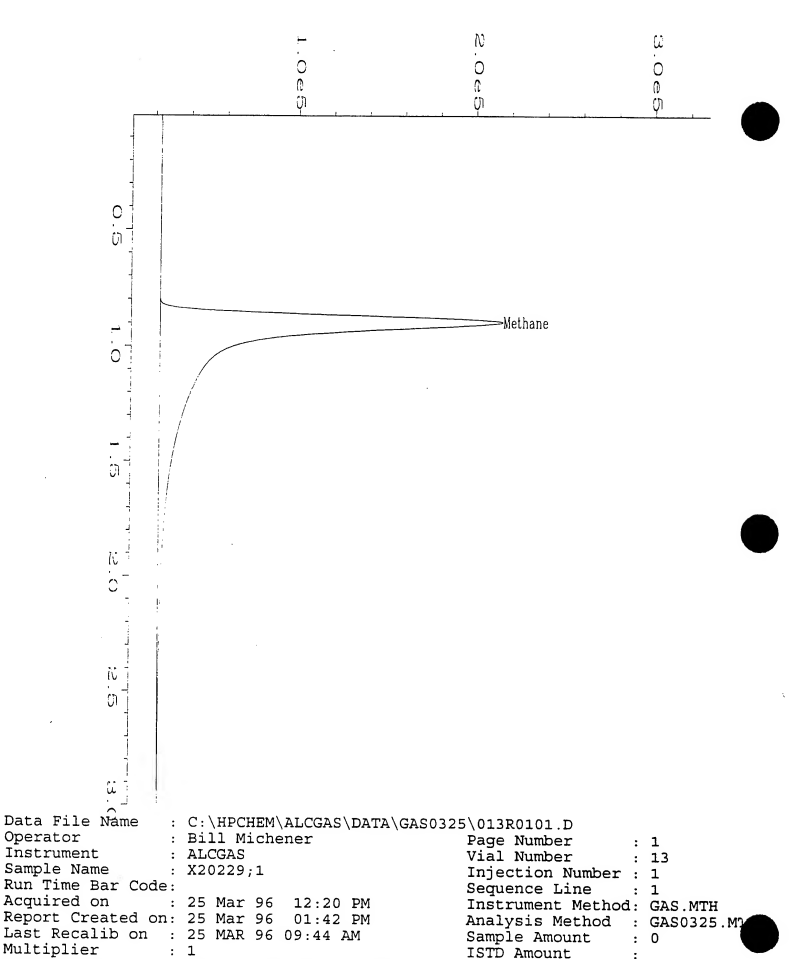
U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Lill M Whenor Analyst



:

Multiplier 1 Sample Info : 96-0741;BE-MP44-01;Water

Methane Report Form

t Sample Number	: BE-MP11-01	Client Project No.	: 722450.14020
Sample Number	: X20228	Lab Project No.	: 96-0741
Date Sampled	: 3/12/96	Dilution Factor	: 2.00
Date Received	: 3/13/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/25/96	Matrix	: Water
Date Analyzed	: 3/25/96	Lab File No.	: GAS0325012

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	0.176	0.004

nperature	:	72.2 F	Saturation	Meth _	0.042503335
	·	0.25 ml	Concentration		
Volume of Sample pace created	:	43 ml	Concentration	Meth	0.133868262
	:	4 ml	in Head Space		
Memáne Area	:	494.197 ug			

Atomic weight(Methane) : ____

<u>16</u> g

QUALIFIERS:

E = Extrapolated value.

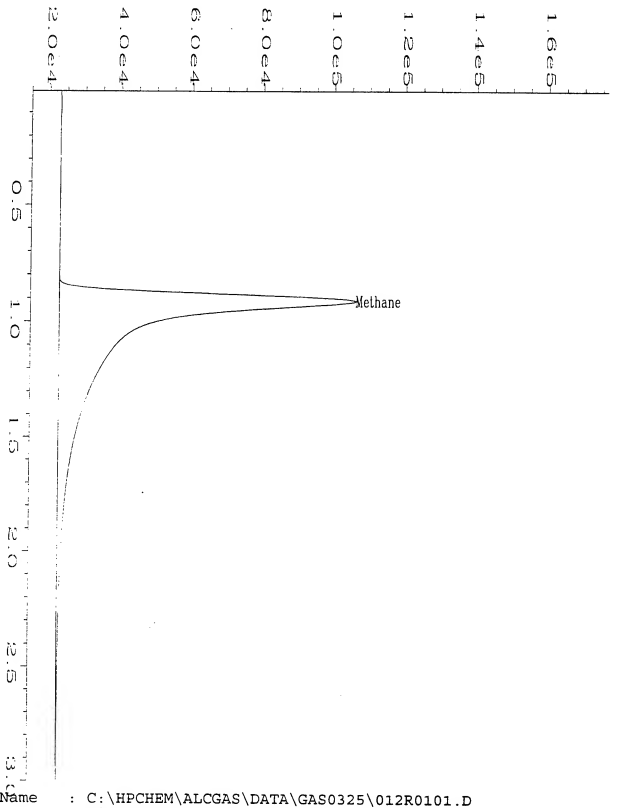
U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Analyst



Data File Name :	C:\HPCHEM\ALCGAS\DATA\GAS032	5\012R0101.D	
Operator :	Bill Michener	Page Number :	1
Instrument :		Vial Number :	
Sample Name :		Injection Number :	
Run Time Bar Code:	•	Sequence Line :	
Acquired on :	25 Mar 96 12:16 PM	Instrument Method:	
Report Created on:	25 Mar 96 01:42 PM	Analysis Method :	
	25 MAR 96 09:44 AM	Sample Amount :	0.0000000000000000000000000000000000000
Multiplier :		ISTD Amount :	ŭ
	06 0540 55 1554	LOID IMMOUIL	

Sample Info : 96-0741;BE-MP11-01;Water

Methane Report Form

Sample Number	: BE-MP42-01	Client Project No.	: 722450.14020
sample Number	: X20227	Lab Project No.	: 96-0741
Date Sampled	: 3/12/96	Dilution Factor	: 20.00
Date Received	: 3/13/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/25/96	Matrix	: Water
Date Analyzed	: 3/25/96	Lab File No.	: GAS0325011

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	0.22	0.04

mperature	:	72.6 F	Saturation	Meth	0.052537778
nount Injected	÷	0.025 ml	Concentration		
Tolume of Sample	:	43 ml	Concentration	Meth	0.165348271
pace created	:	4 ml	in Head Space		
Memane Area	:	61.087 ug			

Atomic weight(Methane) : _____ g

QUALIFIERS:

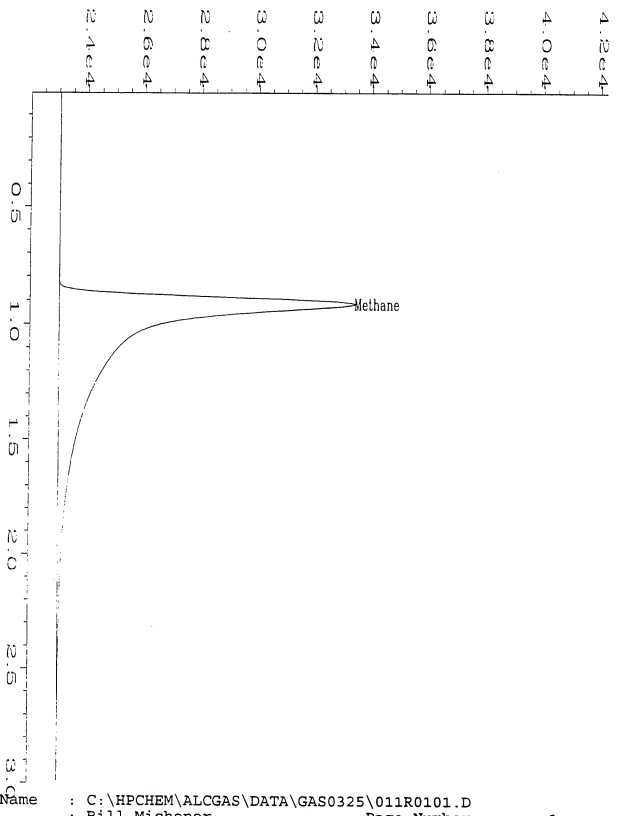
E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.



Data File Name :	C:\HPCHEM\ALCGAS\DATA\GAS032	5\011R0101.D	
Operator :	Bill Michener	Page Number :	1
Instrument :		Vial Number :	
Sample Name :	X20227;20	Injection Number :	
Run Time Bar Code:		Sequence Line :	1
Acquired on :	25 Mar 96 12:04 PM	Instrument Method:	
Report Created on:	25 Mar 96 01:42 PM	Analysis Method :	
Last Recalib on :	25 MAR 96 09:44 AM	Sample Amount :	
Multiplier :	1	ISTD Amount :	,
Sample Info :	96-0741;BE-MP42-01;Water	•	
-			

Methane Report Form

Sample Number	: BE-MP22-01	Client Project No.	: 722450.14020
ample Number	: X20226	Lab Project No.	: 96-0741
Date Sampled	: 3/12/96	Dilution Factor	: 20.00
Date Received	: 3/13/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/25/96	Matrix	: Water
Date Analyzed	: 3/25/96	Lab File No.	: GAS0325010

Compound Name	Sample Cas Number Concentration RL mg/L mg/L			
Methane	74-82-8	0.20	mg/L 0.04	

nperature	:	71.2 F	Saturation	Meth	0.048592736
.iount Injected	:	. 0.025 ml	Concentration		
To Volume of Sample	:	43 ml	Concentration	Meth	0.153335846
pace created	:	4 ml	in Head Space		
Memane Area	:	56.5 ug			

<u>16</u> g

QUALIFIERS:

E = Extrapolated value.

Atomic weight(Methane)

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

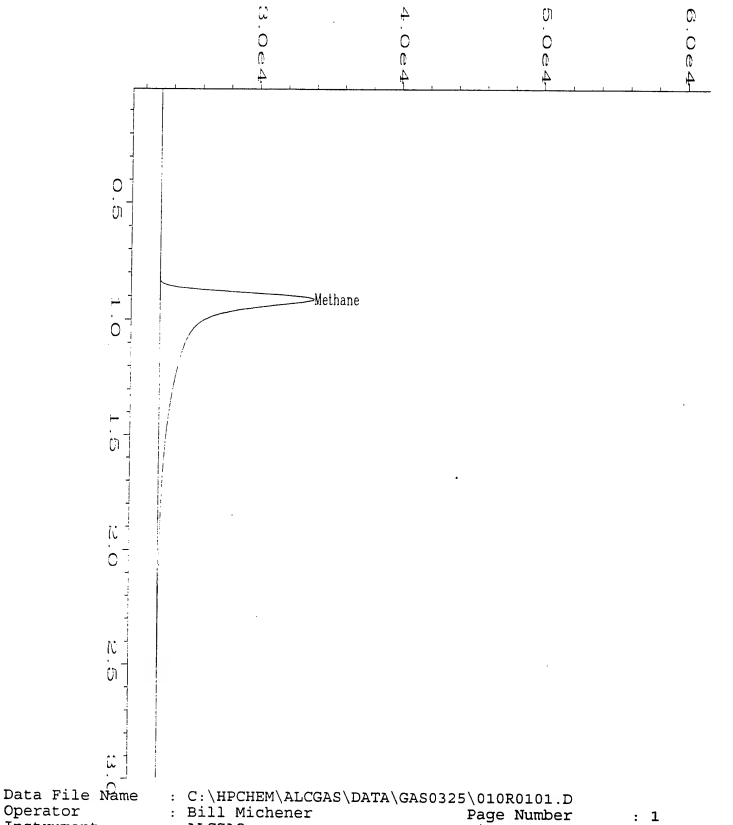
RL = Reporting Limit.

NA = Not Available/Not Applicable.

M Let and Analyst

Approved

AF0741.XLS



Operator Instrument ALCGAS Vial Number 10 Sample Name : X20226;20 Injection Number : 1 Run Time Bar Code: Sequence Line : 1 Acquired on : 25 Mar 96 11:30 AM Instrument Method: GAS.MTH Report Created on: 25 Mar 96 Analysis Method 01:41 PM : GAS0325.M Last Recalib on : 25 MAR 96 09:44 AM Sample Amount : 0 Multiplier ISTD Amount :

: 1 Sample Info : 96-0741;BE-MP22-01;Water

Methane Report Form Method Blank Report

nod Blank Number

: GB032596

: 722450.14020

Date Extracted/Prepared

: 3/25/96

Client Project No. Lab Project No.

: 96-0741

Date Analyzed

: 3/25/96

Dilution Factor

: 1.00

Method

: RSKSOP-175

Matrix

: Water

Lab File No.

: GAS0325002

Sample

Compound Name Cas Number Concentration RL mg/L mg/L

Methane

74-82-8

U

0.002

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

AF0741.XLS

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Data File Name
                     : C:\HPCHEM\ALCGAS\DATA\GAS0325\002R0101.D
                     : Bill Michener
                                                          Page Number
                                                                               : 1
Instrument
                     : ALCGAS
                                                          Vial Number
Sample Name
                     : GB032596
                                                          Injection Number: 1
Run Time Bar Code:
                                                          Sequence Line
                                                                             : 1
Acquired on
                    : 25 Mar 96
                                    08:51 AM
                                                          Instrument Method: GAS.MTH
Report Created on: 25 Mar 96
                                    09:45 AM
                                                          Analysis Method : GAS0325.N
```

Displaced 4ml of deionized water in 43ml vial with Helium, shook for 5 min and injected 500ml

Sample Amount

ISTD Amount

: 0

Operator

Multiplier

Sample Info

Last Recalib on : 25 MAR 96 09:44 AM

: Gas Method Blank

Date & Time Rec'd: 3/13/96 09 5 Shipped Via: For	115409095
(Airbil	1 # if applicable)
Client: (Sons t)	
Client Project ID(s):	
EAL Project $\#(s): 96-0+1$ EAL Cooler	(s): (?) N
Cooler#	
Ice packs P'N Y N Y N Y N	X N
Temperature °C ()	
La Mord on temp blank (2)	N N/A
1. Custody seal(s) present: Seals on cooler intact Seals on bottle intact	
2. Chain of Custody present:	
3. Samples Radioactive: (Comment on COC if > 0.5mr/h)	<u> </u>
4. Containers broken or leaking: (Comment on COC FY)	
5. Containers labeled:	
6. CCC agrees w/ bottles received: (Comment on COC IF)	
7. CCC agrees w/ labels: (Comment on COC if N)	. ———
8. Headspace in vials-waters only: (Comment on COCIFY)	
9. VCA samples preserved:	
10. pH measured on metals, cyanide or phenolics*: List discrepancies *Non-EAL provided containers only, water samples only	
11. Metal samples present: Total , Dissolved , TCLP	X
D or PD to be filtered:	
T,TR,D,PD to be Preserved:	
12. Short holding times: Specify parameters NO2/NO3	
13. Multi-phase sample(s) present:	<u></u>
14. COC signed w/ date/time:	-
Comments:	
	·
(Additional comments on back) WWW Z/12/6/	
Custodian Signature/Date:	

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Total Organic Carbon

Date Sampled Date Received	: 3/12/96 : 3/13/96	Client Project ID. Lab Project Number		722450.14020
Date Prepared	: 3/14/96	Method		EPA 415.1
Date Analyzed	: 3/14/96	Detection Limit	:	1.0 mg C/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	TOC	mg C/L	Dilution <u>Factor</u>
X20226	BE-MP22-01	Water	4.3		1
X20227	BE-MP42-01	Water	4.6		1
X20228	BE-MP11-01	Water	2.1		1
X20229	BE-MP44-01	Water	2.2		1

Method Blank

(3/14/96)

<1.0

Quality Assurance

		Spike Amount (mgC/L)	Sample Result (mgC/L)	Spike Result (mgC/L)	% Recovery
X20226	BE-MP22-01 Matrix Spike	10.0	4.3	15.2	109
X20226	BE-MP22-01 Matrix Spike Du	10.0	4.3	14.5	102
MS/MSD I	RPD				6.2

Analyst

Evergreen Analytical Sample Log She	et Pro	oject #	# <u>96-07</u>	41
Date(s) Sampled: 03/12/96 COC	Date	Due:	03/27/	96
te Received: 03/13/96 0915	Holding Time	(s):	03/14-	·NO ₂ , NO ₂ ,
Client Project I.D. 722450.14020		3/26-	ALKALI STANDA	NITY
Client: PARSONS ENGINEERING SCIEN	CE, INC. Cooler R	eturn	N/A	
Address: 1301 MARINA VILLAGE PKWY				
ALAMEDA, CA 94501				4090954
Contact: MICHAEL PHELPS				
Phone #510-769-0100 Fax #510-769				
Special Invoicing/Billing				
Special Instructions				
Lab Client				
	Analysis	Mtx	Btl	Loc
X20226A BE-MP22-01	TOC	W	125A	CL4
X20227A BE-MP42-01	TOC	W	125A	
X20228A BE-MP11-01	TOC	W	125A	
¥2 <u>0229A BE-MP44-01</u>	TOC	W	125A	
226B BE-MP22-01 NO ₂	,NO ₃ ,SO ₄ ,Cl ⁻	W	125P	CL4
X20227B BE-MP42-01 NO2	, NO ₃ , SO ₄ , Cl ⁻	W	125P	CL4
X 8B BE-MP11-01 NO2	, NO ₃ , SO ₄ , Cl ⁻	W	125P	CL4
X20229B BE-MP44-01 NO ₂	, NO ₃ , SO ₄ , Cl	W	125P	CL4
X20226C BE-MP22-01	MMONIA	W	500P	CL4
X20227C BE-MP42-01	MMONIA	W	500P	CL4
X20228C BE-MP11-01	MMONIA	W	500P	CL4
X20229C BE-MP44-01	MMONIA	W	500P	CL4
X20226D BE-MP22-01 A	LKALINITY	W	125P	CL4
X20227D BE-MP42-01	LKALINITY	W	125P	CL4
	LKALINITY	W	125P	CL4
	LKALINITY	W	125P	CL4
	ETHANE	W	40V	2
	ETHANE	W	40V	2
***************************************	ETHANE	W	40V	2
,	ETHANE	W	40V	2
R=Sample to be returned				
.te GC/MS GC X_ Metals	Wet Chem X_	HPLC	_	SxPrep
SxRec C QA/QC	<u>C</u> Acctg <u>C</u> Fil	e <u>c</u>	rig	
Page 1 of 1 Page(s)			/Date	: MM 3/17/96

CHAIN OF CUSTODY RECORD / ANALYTICAL SERVICES REQUEST

CLIENT CONTACT (print) MICLAGE PLOS PROJECT I.D. EAL. QUOTE # P.O.# TURNAROUND REQUIRED* \$\superigoning{\text{TED}}\$ \text{Other (Specify)} \\ *expedited turnaround subject to additional fee Do not write Do not write	EAL Project # 96-074 Custodian WW	20226	22	29				Container Size	
PO.# BY STD (2 wks) Control (Specify) Roadditional fee EAL Do I	m 1 0		-					3 5	
CLIENT CONTACT (print) MICMAE PROJECT I.D. EAL. QUOTE # P.O.# TURNAROUND REQUIRED* \$\square\$ STD (2 wks) Cother (Specificated turnaround subject to additional fee	All and								
TACT	Methons	×	<u> </u>	X			1		}
CLIENT CONTA	11550 Ned AWMONIA	X	y y	X	11		1		4
PROJE SAL. O URNA EXPEDI	Tain Parally SHOILD	>	XX	X X	+	+			
CLI CLI CLI EAU EAU TUF 'ext	Oissolved Metals below) Oissolved Metals below) Oissolved Metals below)		7-		+				
igo U	Total Metals-DW / NPDES / SW846 (Circle & list metals below) Dissolved Metals - DW / SW846 (circle & list matals - DW / SW846)]
Analytical Inc. 4036 Younglield St. 4036 Younglield St. 6303) 425-6021 FAX (303) 425-6854 (800) 845-7400 FAX RESULTS Y / N ANALYSIS RE	TEPH 8015mod. (Gasoline)	\sqcup					1-1-	_	
Iytical Inc ingled St. idge, Colorado 8 5-6021 5-7400 ESULTS Y /	S IIO/I:o	H			1 1		-	+	-
ytica grield gge, Co 6021 7400 3ULTS	BTEX 8020/602 (circle)/MTPC 4							+-	1
Evergreen Analytical Inc. 4036 Younglield St. 4036 Younglield St. Wheat Ridge, Colorado 80 (303) 425-6021 FAX (303) 425-6854 (800) 845-7400 FAX RESULTS Y / N ANALYSIS F	PCB Screen								
4030 4030 Whe (303) FAX (800) FAX	209/0808 SBO-100								
gree	Pesticides 8080/608 (circle)					+		╁	
Ever	1999	V;	XX	X			P	3	
0 0	TCLP VOA/BNA/Pest/Herb/Metals (circle) VOA 8260/624/524.2 (circle)								1
36							\vdash	-	
ASE PEWT STECTO 94501 FAX# STO 769	biloS \ liO				1	++	+		
WA S	Water-Drinking/Discharge/Ground (circle)	٧:	××	X.					
GUSO GUSO FAX #	No. of Containers	4	7+	4					
o do	TIME	07	1130 1330	345					
				ऽभ्हा त्र है।	+	+			
S ES CA CA Oloo	TINT ation:	5/2	96/21/ 96/21/	<u>ध</u> ्रीया					
Mavium STATE CA STATE CA 69 010	ation D, SAN,	3/12/	3/1/2	12					
COMPANY Paysons ES ADDRESSIZOI Maying CITY Alawerla STATE CA PHONE SID 769 0100 Sampler Name; (signature) Lawer Lynn (print) favie Torigin Michael Calabara	Evergreen Analytical Cooler No. Cooler Received Please PRIN Please PRIN all information: CLIENT SAMPLE DATE DATE	8	2 -	10					
Par 301	green Analytical Cook ler Received Please Pall info CLIENT SAMPLE IDENTIFICATION	BE-MP22-01	BE-MP42-0	BE-MP4-0					
SI3	Please Please al CLIENT SAMPLE	S S	N P P	25					ions:
COMPANY POC ADDRESS 13CH CITY ARAMERIC PHONE SAMPIER NAME; (Signature) Lama (print) Lavice 16	Evergreen Analyt Cooler Received Ple CLIER SAMP	ريا	1 2 2	1					Instructions:
COM ADE CIT PHC Sar (sign	Coo	Ø 6	かな	9			<u> </u>	8	Ĕ

lature) Relinqui sature

| 3/10 Date/Time | Received by: (Signature)

inquished by: (Signature)

Date/Time Regel

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Sulfate</u> mg/L	Dilution <u>Factor</u>
X20031	BE-MP15-01	Water	11.4	1
X20032	BE-MP17-01	Water	3.4	1
X20033	BE-MP-8-01	Water	467	50
X20034	BE-MP-41-01	Water	458	50

Method Blank (3/12/96)

< 0.25

Quality Assurance

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20032	BE-MP17-01 Matrix Spike	10.0	3.4	13.0	95
X20032	BE-MP17-01 Matrix Spike Du	p 10.0	3.4	12.3	89
MS/MSD	RPD				7.0

Analyst

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled : 3/11/96 Date Received : 3/12/96 Date Prepared : 3/12/96 Date Analyzed : 3/12/96	Lab Project Number Method	:	722450.14020 96-0706 EPA 300.0 0.076 mg/L
---	------------------------------	---	--

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Nitrite-N</u> mg/L	Dilution <u>Factor</u>
X20031	BE-MP15-01	Water	<0.076	1
X20032	BE-MP17-01	Water	<0.076	1
X20033	BE-MP-8-01	Water	<0.76**	10
X20034	BE-MP-41-01	Water	<0.76**	10

Method Blank (3/12/96)

< 0.076

Quality Assurance *

	:	Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20032	BE-MP17-01				
	Matrix Spike	10.0	<0.25	9.7	97
X20032	BE-MP17-01				
	Matrix Spike Du	p 10.0	<0.25	8.9	89
MS/MSD F	RPD				8.0

Quality assurance results reported as Nitrite (NO₂).

** = Increased detection limit due to Matrix Interference.

st Appr

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled	: 3/11/96	Client Project ID.	:	722450.14020
Date Received	: 3/12/96	Lab Project Number	:	96-0706
Date Prepared	: 3/12/96	Method	:	EPA 300.0
Date Analyzed	: 3/12/96	Detection Limit	:	0.056 mg/L

Evergreen Sample #	Client <u>Sample ID.</u>	<u>Matrix</u>	Nitrate-N mg/L	Dilution <u>Factor</u>
X20031	BE-MP15-01	Water	0.18	1
X20032	BE-MP17-01	Water	2.5	1
X20033	BE-MP-8-01	Water	148	50
X20034	BE-MP-41-01	Water	141	50

Method Blank (3/12/96)

< 0.056

Quality Assurance *

	<u> </u>	Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20032	BE-MP17-01 Matrix Spike	10.0	11.2	21.4	102
X20032	BE-MP17-01 Matrix Spike Du	o 10.0	11.2	21.0	98
MS/MSD	RPD				4.0

^{• =} Quality assurance results reported as Nitrate (NO₃).

Analyst

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

Date Sampled : 3/11/96 Date Received : 3/12/96

Client Project ID. : 722450.14020 Lab Project Number : 96-0706

Date Received : 3/12/96
Date Prepared : 3/18/96
Date Analyzed : 3/18/96

Method : Standard Method 4500-NH₃F

Detection Limit : 0.80 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Ammonia-N	Dilution <u>Factor</u>
X20031	BE-MP15-01	Water	<0.80 mg/L	1
X20032	BE-MP17-01	Water	<0.80 mg/L	1
X20033	BE-MP-8-01	Water	3.3 mg/L	1
X20034	BE-MP-41-01	Water	1.9 mg/L	1

Method Blank

(3/18/96)

<0.80 mg/L

Quality Assurance

	3	Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20031	BE-MP15-01 Matrix Spike	10.0	<0.80	9.5	95
X20031	BE-MP15-01 Matrix Spike Du	10.0 p	<0.80	9.4	94
MS/MSD E	2PD				

MS/MSD RPD

1.2

Analyst Hole

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

 Date Sampled
 : 3/11/96
 Client Project ID.
 : 722450.14020

 Date Received
 : 3/12/96
 Lab Project Number
 : 96-0706

 Date Prepared
 : 3/12/96
 Method
 : EPA 300.0

 Date Analyzed
 : 3/12/96
 Detection Limit
 : 0.25 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Chloride</u> mg/L	Dilution Factor
X20031	BE-MP15-01	Water	84.4	10
X20032	BE-MP17-01	Water	20.3	1
X20033	BE-MP-8-01	Water	925	50
X20034	BE-MP-41-01	Water	885	50

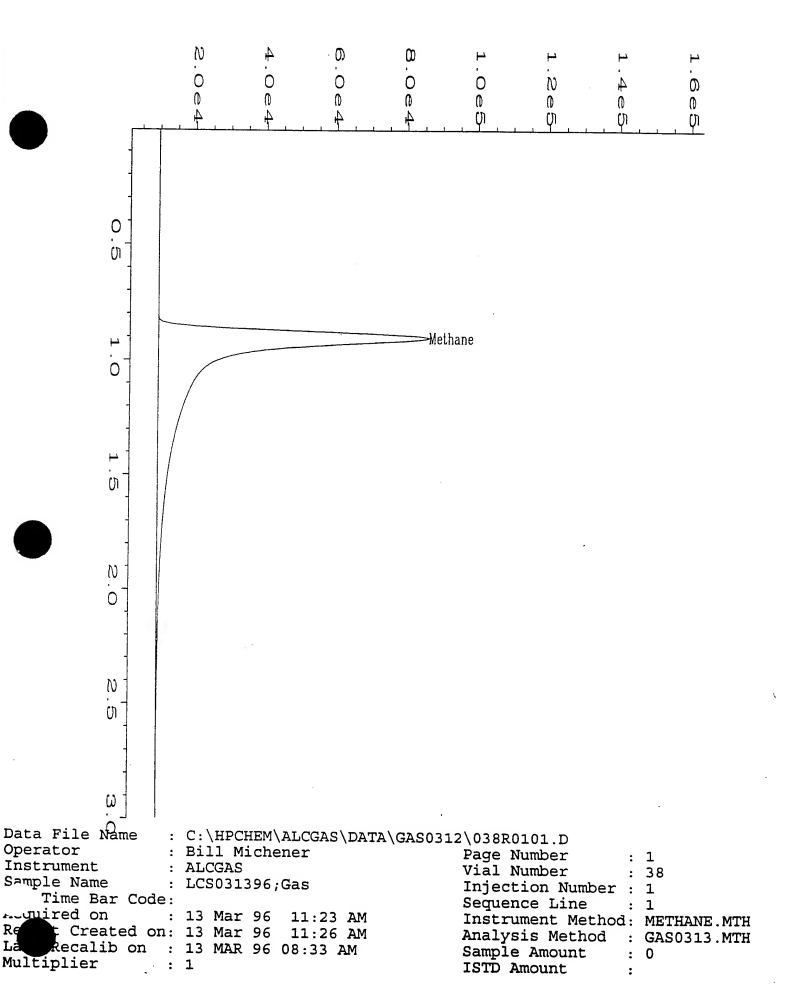
Method Blank (3/12/96)

< 0.25

Quality Assurance

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X20032	BE-MP17-01 Matrix Spike	10.0	20.3	29.7	94
X20032	BE-MP17-01 Matrix Spike Du	up 10.0	20.3	30.0	97
MS/MSD F	RPD .				3.3

Analyst



4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

Date Sampled Date Received Date Prepared	: 3/11/96 : 3/12/96 : 3/18/96	Client Project ID. Lab Project Number Method	: EPA 310.1
Date Analyzed	: 3/18/96	Detection Limit	: 5.0 mg CaCO ₃ /L

Client <u>Sample ID.</u>	<u>Matrix</u>	Total <u>Alkalinity</u> (mg CaCO ₃ /L)	Dilution <u>Factor</u>
BE-MP17-01	Water	68.7	1
BE-MP-8-01	Water	1220	1
BE-MP-41-01	Water	1250	1
	Sample ID. BE-MP17-01 BE-MP-8-01	Sample ID. Matrix BE-MP17-01 Water BE-MP-8-01 Water	Sample ID. Matrix Alkalinity (mg CaCO ₃ /L) BE-MP17-01 Water 68.7 BE-MP-8-01 Water 1220

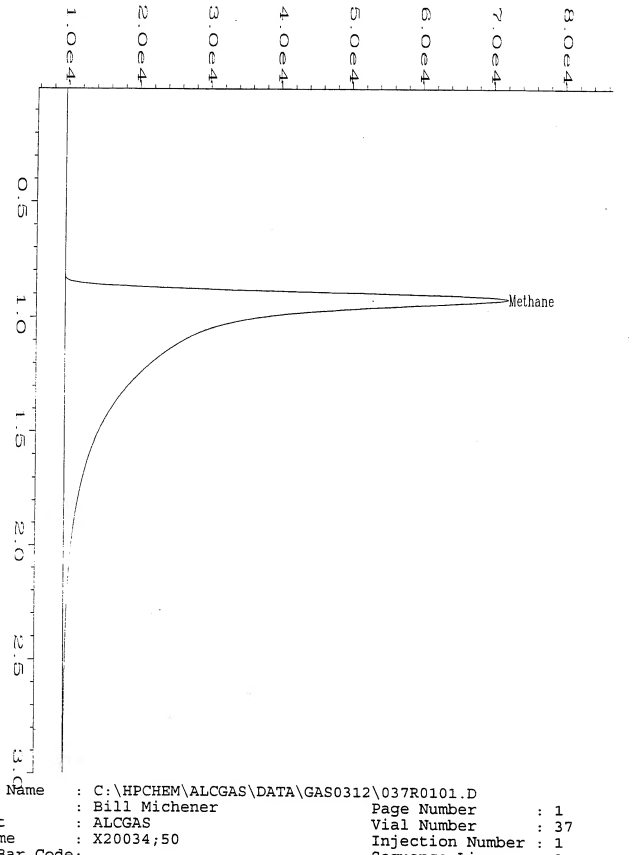
Method Blank (3/18/96)

< 5.0

Quality Assurance

116	97
	116

Haliman Haliman



Data File Name :	C:\HPCHEM\ALCGAS\DATA\GAS03	12\037R0101.D	
Operator :	Bill Michener	Page Number :	1 .
Instrument :	ALCGAS	Vial Number :	
Sample Name :	X20034;50	Injection Number :	
Time Bar Code:		Sequence Line :	
_uired on :	13 Mar 96 11:14 AM	Instrument Method:	
ke t Created on:	13 Mar 96 11:21 AM	Analysis Method :	
L Recalib on :	13 MAR 96 08:33 AM	Sample Amount :	0
	1	ISTD Amount	
Sample Info :	96-0706;BE-MP-41-01:Water	•	

Evergreen Analytical, Inc. 4036 Youngfield, Wheat Ridge, CO 80033 (303) 425-6021

RSK-175 Gas Method Methane LCS Report Form

LCS No.

: LCS031396

EPA Method No.

: RSKSOP-175

Date Prepared

: 3/13/96

Matrix

: Water

Date Analyzed

: 3/13/96

Method Blank

: GB031396

E.A. LCS Source No.

: 1723

Lab File No.

: GAS0312038

Compound	Spike Added (ug)	Method Blank Concentration (ug)	LCS Concentration (ug)	LCS %REC	QC Limits %REC
Methane Gas	500	. 0	361	72	67-85

Spike Recovery:

	\sim
	11

out of (1) outside limits.

Note: The LCS was made by taking the sample and displacing 4ml of headspace with a 1% methane gas and shaking the VOA for 5 minutes. Then injecting 50 ul from the headspace into the GC resulting in a theoretical concentration of 500 ug.

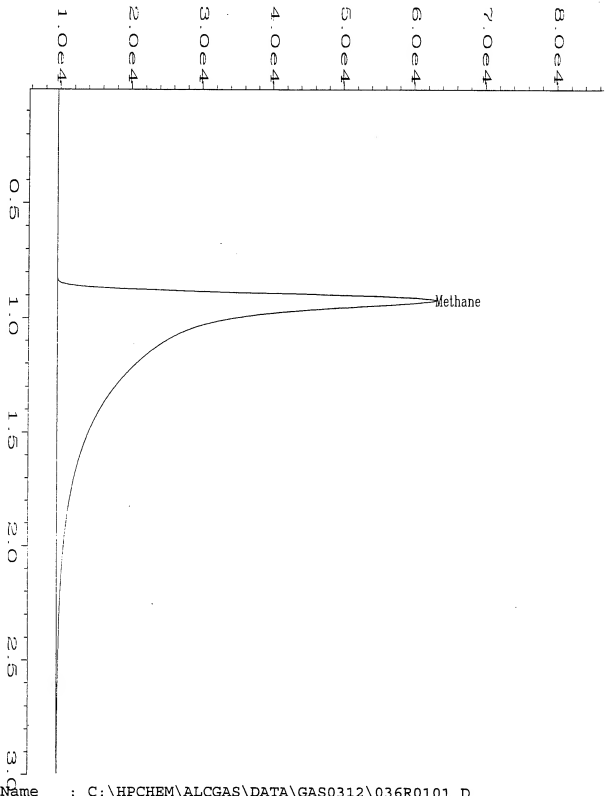
NOTES:

* = Values outside of QC limits. NA = Not analyzed/not available.

Analyst

Approved

LCS0313.XLS; 3/13/96



Data File Name : C:\HPCHEM\ALCGAS\DATA\GAS0312\036R0101.D Operator : Bill Michener Page Number : 1 Instrument : ALCGAS Vial Number : 36 Sample Name : X20033;50 Injection Number: 1 Time Bar Code: Sequence Line : 1 mired on : 13 Mar 96 11:10 AM Instrument Method: METHANE.MTH t Created on: 13 Mar 96 11:21 AM Analysis Method : GAS0313.MTH

Recalib on : 13 MAR 96 08:33 AM Sample Amount : 0 Multiplier : 1 ISTD Amount :

Sample Info : 96-0706; BE-MP8-01; Water

EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP-41-01	Client Project No.	: 722450.14020
Lab Sample Number	: X20034	Lab Project No.	: 96-0706
Date Sampled	: 3/11/96	Dilution Factor	: 50.00
Date Received	: 3/12/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/13/96	Matrix	: Water
Date Analyzed	: 3/13/96	Lab File No.	: GAS0312037

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	3.9	0.1

Temperature	:	72.2 F	Saturation	Meth	0.946625
Amount Injected	:	0.01 ml	Concentration		
Total Volume of Sample	:	43 ml	Concentration	Meth	2.98149
Head space created	:	4 ml	in Head Space		
Methane Area	:	440.268 ug			***

Atomic weight(Methane) : _____ g

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

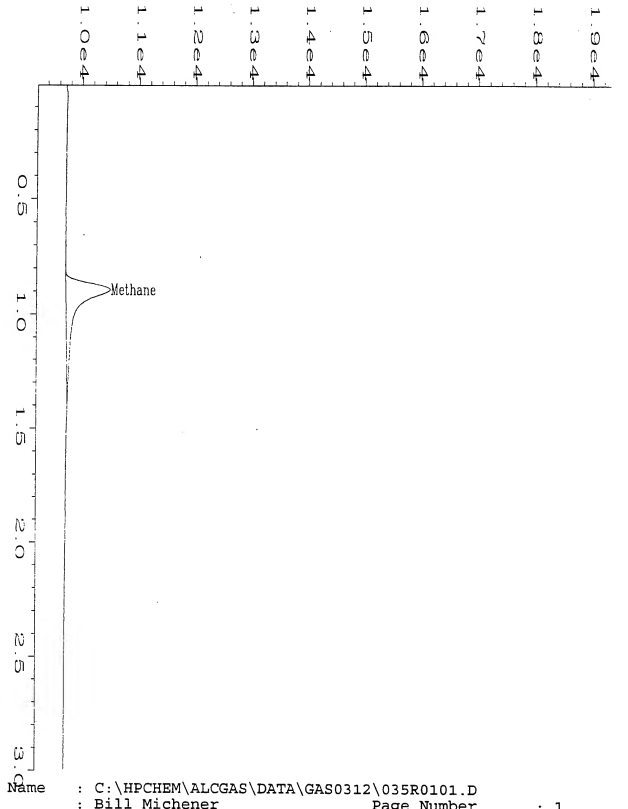
B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Analyst Analyst

K. Hollman
Approved



Data File Name Operator : Bill Michener Page Number : 1 Instrument : ALCGAS Vial Number : 35 Sample Name : X20032Dup;1 Injection Number: 1 Time Bar Code: Sequence Line wired on : 13 Mar 96 10:58 AM Instrument Method: METHANE.MTH t Created on: 13 Mar 96 11:21 AM Analysis Method : GAS0313.MTH

Recalib on : 13 MAR 96 08:33 AM Sample Amount : Multiplier : 1 ISTD Amount :

Sample Info : 96-0706;BE-MP17-01;Water

EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP8-01	Client Project No.	: 722450.14020
Lab Sample Number	: X20033	Lab Project No.	: 96-0706
Date Sampled	: 3/11/96	Dilution Factor	: 50.00
Date Received	: 3/12/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/13/96	Matrix	: Water
Date Analyzed	: 3/13/96	Lab File No.	: GAS0312036

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	3.5	0.1

Temperature	:	72.8 F	Saturation	Meth	0.84551F '
Amount Injected	:	0.01 ml	Concentration		-
Total Volume of Sample	:	43 ml	Concentration	Meth	2.66002
Head space created	;	4 ml	in Head Space		
Methane Area	:	393.241 ug			

Atomic weight(Methane) : 16 g

QUALIFIERS:

E = Extrapolated value.

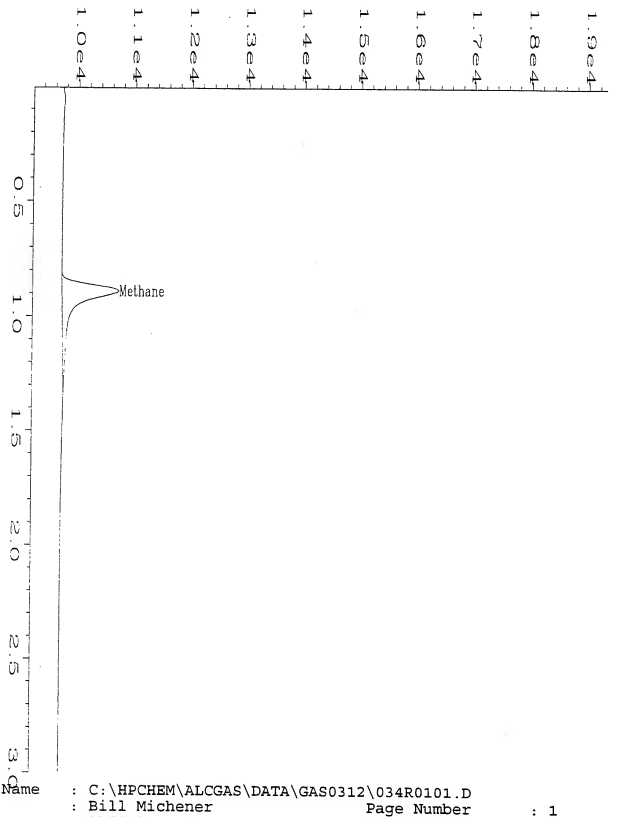
U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Analyst



Data File Name	:	C:\HPCHEM\ALCGAS'	\DATA\GAS0312\(034R0101.D		
Operator	:	Bill Michener				1
Instrument	:	ALCGAS		ial Number	-	_
Sample Name	:	X20032;1		njection Number		
Time Bar Code	:	·	Se	equence Line		1
uired on	:	13 Mar 96 10:54	AM II	nstrument Method:		

THANE.MTH Recalib on: 13 Mar 96 11:21 AM Recalib on: 13 MAR 96 08:33 AM Multiplier: 1
Sample Info: 96-0706;BE-MP17-01;W Analysis Method Sample Amount ISTD Amount : GAS0313.MTH

: 0 : : 1 : 96-0706;BE-MP17-01;Water

EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP17-01	Client Project No.	: 722450.14020
Lab Sample Number	: X20032Dup	Lab Project No.	: 96-0706
Date Sampled	: 3/11/96	Dilution Factor	: 1.00
Date Received	: 3/12/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/13/96	Matrix	: Water
Date Analyzed	: 3/13/96	Lab File No.	: GAS0312035

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

Temperature	:	74.5 F	Saturation	Meth	0.00013F1 3
Amount Injected	:	0.5 ml	Concentration		-
Total Volume of Sample	:	43 ml	Concentration	Meth	0.000424
Head space created	•	4 ml	in Head Space		
Methane Area	:	3.145 ug			

Atomic weight(Methane) : _____ 16 g

QUALIFIERS:

E = Extrapolated value.

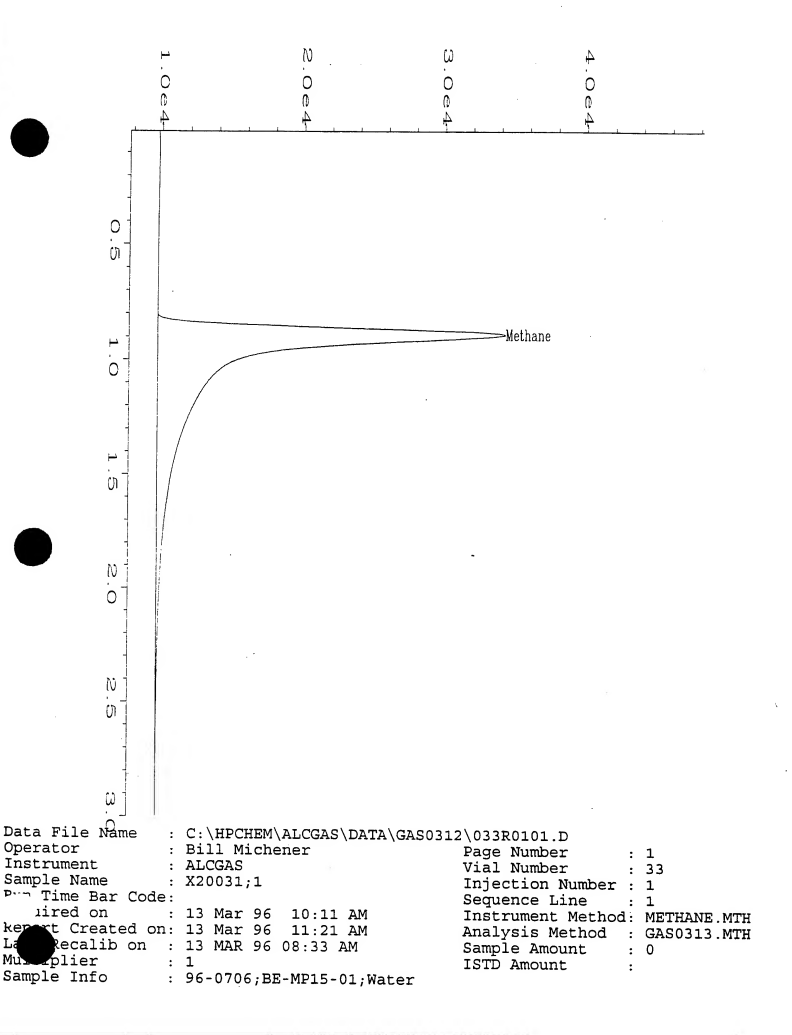
U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Analyst



EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP17-01	Client Project No.	: 722450.14020
Lab Sample Number	: X20032	Lab Project No.	: 96-0706
Date Sampled	: 3/11/96	Dilution Factor	: 1.00
Date Received	: 3/12/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/13/96	Matrix	: Water
Date Analyzed	: 3/13/96	Lab File No.	: GAS0312034

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

Temperature	:	74.3 F	Saturation	Meth	0.00014/
Amount Injected	:	0.5 ml	Concentration		
Total Volume of Sample	:	43 ml	Concentration	Meth	0.000453
Head space created		4 ml	in Head Space		
Methane Area		3 365 ug			

Atomic weight(Methane) :16	Atomic weight(Methane)	:	16	g
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QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

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Data File Name
                      : C:\HPCHEM\ALCGAS\DATA\GAS0312\022R0101.D
                      : Bill Michener
                                                             Page Number
                                                                                   : 1
Instrument
                      : ALCGAS
                                                             Vial Number
                                                                                   : 22
```

Sample Name : GB031396 Injection Number: 1 Time Bar Code: Sequence Line : 1 uired on : 13 Mar 96 08:45 AM Instrument Method: METHANE.MTH Re Created on: 13 Mar 96 11:20 AM Analysis Method : GAS0313.MTH : 13 MAR 96 08:33 AM Recalib on Sample Amount : 0 Multiplier : 1 ISTD Amount : Sample Info : Gas Method Blank

Displaced 4ml of deionized water in 43ml vial with Helium, shook for 5 min and injected 500ml

Operator

EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP15-01	Client Project No.	: 722450.14020
Lab Sample Number	: X20031	Lab Project No.	: 96-0706
Date Sampled	: 3/11/96	Dilution Factor	: 1.00
Date Received	: 3/12/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/13/96	Matrix	: Water
Date Analyzed	: 3/13/96	Lab File No.	: GAS0312033

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	0.023	0.002

Temperature	:	74.3 F	Saturation	Meth	0.00548'
Amount Injected	:	0.5 ml	Concentration		
Total Volume of Sample	:	43 ml	Concentration	Meth	0.01721
Head space created	:	4 ml	in Head Space		
Methane Area	:	127.573 ug			

Atomic weight(Methane) : _____ 16 g

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

K Hollman Approved

Anaiyst

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Total Organic Carbon

Date Sampled	: 3/11/96	Client Project ID.	:	722450.14020
Date Received	: 3/12/96	Lab Project Number	:	96-0706
Date Prepared	: 3/14/96	Method	:	EPA 415.1
Date Analyzed	: 3/14/96	Detection Limit	:	1.0 mg C/L

Evergreen Sample #	Client <u>Sample ID.</u>	<u>Matrix</u>	TOC mg C/L	Dilution <u>Factor</u>
X20031	BE-MP15-01	Water	6.6	1
X20032	BE-MP17-01	Water	1.5	1 == ==================================
X20033	BE-MP-8-01	Water	102	1 1500
X20034	BE-MP-41-01	Water	103	1

Method Blank

(3/14/96)

<1.0

Quality Assurance

		Spike Amount (mgC/L)	Sample Result (mgC/L)	Spike Result (mgC/L)	% Recovery
X20031	BE-MP15-01 Matrix Spike	10.0	6.6	16.5	99
X20031	BE-MP15-01 Matrix Spike Du	10.0 P	6.6	17.0	104
MS/MSD F	RPD				4.8

EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form Method Blank Report

Method Blank Number Date Extracted/Prepared

Date Analyzed

: GB031396

: 3/13/96 : 3/13/96

Client Project No.

: 722450.14020

Lab Project No.

: 96-0706

Dilution Factor Method

: 1.00 : RSKSOP-175

Matrix

: Water

Lab File No.

: GAS0312022

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

CHAIN OF CUSTODY RECORD		ALYTICAL SERVICES REQUEST	
	Evergreen Analytical Inc.		Page of
COMPANY THESE S	4036 Youngfield St.	CLIENT CONTACT (print) MICHAREL PHELDS	MEL PHELPS
ADDRESS [30] MAKINA VILLAGE PLUY ST 200	Wheat Hidge, Colorado 80033 (303) 425-6021	PROJECT I.D.	
CITY & AMEDA STATE CA ZIP Q 450/	FAX (303) 425-6854 (800) 845-7400	EAL. QUOTE #	P.O.#
PHONE \$10 764 0100 1 FAX # 510 769 9244	FAX RESULTS Y / N	TURNAROUND REQUIRED* STD (2 wks) STD UST (3 day)	(2 wks) STD UST (3 day)
Sampler Name:		Other	Other (Specify)
(signature) i'milli M much pulle Miller		expedited turnaround subject to additional fee	nal fee
(print) Proflage H. Fright	ANALYSIS REQUESTED, \$	STED.	EAL use only
TI NOW BUT HERE		स्र	Do not write
Evergreen Analytical Cooler No.	(6 (e)	では	in shaded area

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Please

Cooler Received Cooler Received Please PRINT Please all information: CLIENT SAMPLE DATE IDENTIFICATION SAMPLED TIME	bilo2 \ lio2 egbul2 \ liO	TCLP VOA/BNA/Pest/Herb/Metals (circle) VOA 8260/624/524.2 (circle)	Pest/PCBs 8080/608 (circle)	PCB Screen	BTEX 8020/602 (circle)/MTBE (circle) TRPH 418.1/Oil & Grease 413.1 (circle) TYPH 8015mod. (Gasoline)	Otal Metals-DW / WDDEG	Circle & list metals below) Circle & list metals below) Circle & list metals below) Circle & list metals below)	LICHLINATY SESSIVED ANMOUNT AND CHORIDE AND AND AND AND AND AND AND AND AND AND	ZEIH₩/E	EAL Project #Custodian
BE. WP 15 \$1 3/11 /16 1040 6 X		┥—				4	12	NX NX	+	CAL Sample
X F 081 1911/8 12-FIAM-28		X					×	XXX		
X 7 1181 3/11/96 1515 2 X		X					X	スペス		
(MAE) 8E-MP-8017										
88-M-41-01 3/11/46 15:20 7 X		メ					¥	メスドア		
H.										Location
DD:										Container Size
Instructions:										

Time Received by: (Signature)

Date/Time Relinquished by: (Signature)

Evergreen Analytical Sample Receipt/Check-in Record
Date & Time Rec'd: 3-12-96 0930 Shipped Via: Fix FX 115409273
Client: YARsons ES. (Airbill # if applicable)
Client Project ID(s):
EAL Project #(s):96- EAL Cooler(s): Y
Cooler# 656
Ice packs (Y) N Y N Y N Y N
Temperature °C
1. Custody seal(s) present: Seals on cooler intact Seals on bottle intact
2. Chain of Custody present:
3. Samples Radioactive: (Comment on COC if > 0.5mr/b)
4. Containers broken or leaking: (Comment on COC IF Y)
5. Containers labeled:
6. COC agrees w/ bottles received: (Comment on COC if N)
7. COC agrees w/ labels: (Comment on COC if N)
8. Headspace in vials-waters only: (Comment on COC IF Y)
9. VOA samples preserved:
10. pH measured on metals, cyanide or phenolics*:
11. Metal samples present:
Total, Dissolved, TCLP D or PD to be filtered:
T,TR,D,PD to be Preserved:
12. Short holding times: NO, NO, ———————————————————————————————
13. Multi-phase sample(s) present:
14. COC signed w/ date/time:
Comments:
(Additional comments on back)
Custodian Signature/Date: / / //ast/ >-/2

Evergreen Analytical Sample Log Sheet			Project # <u>96-07</u>	06
Date(s) Sampled: 03/11/96 COC			Date Due: 03/26	'96
te Recei	ived: <u>03/12/96</u> 0	930 Hol	ding Time(s): 03/13-	-NO ₂ , NO ₂ ,
	oject I.D. 7224		3/25-ALKALI Rush STANDA	NITY
Client: D	PCONC PUCTNEEDT	NC COLENOR TWO		
	301 MARINA VILL	NG SCIENCE, INC.	Cooler Return N/A	
	LAMEDA, CA 945		E.A. Cooler # N/A	
	IICHAEL PHELPS	01	Airbill # FEDEX 115 Client P.O	
	-769-0100 Fax	#510-769-92 <i>44</i>	CITERC P.O.	
special In	voicing/Billing			
Special In	structions		····	
Lab	Client			
ID #	ID#	Analysis	Mtx Btl	Loc
X20031A	BE-MP15-01	TOC	W 125A	C4
X20032A	BE-MP17-01	TOC	W 125A	
X20033A	BE-MP-8-01	TOC	W 125A	C4
X20034A	BE-MP-41-01	TOC	W 125A	
1031B	BE-MP15-01	NO_2 , NO_3 , SO_4 , $C1^-$	W 125P	C4
X20032B	BE-MP17-01	NO ₂ , NO ₃ , SO ₄ , Cl ⁻	W 125P	C4
33B	BE-MP-8-01	NO2, NO3, SO4, C1	W 125P	C4
X20034B	BE-MP-41-01	NO ₂ , NO ₃ , SO ₄ , C1	W 125P	C4
X20031C	BE-MP15-01	AMMONIA	W 500P	C4
X20032C	BE-MP17-01	AMMONIA	W 500P	C4
X20033C	BE-MP-8-01	AMMONIA	W 500P	C4
X20034C	BE-MP-41-01	AMMONIA	W 500P	C4
X20032D	BE-MP17-01	ALKALINITY	W 125P	C4
X20033D	BE-MP-8-01	ALKALINITY	W 125P	C4
X20034D	BE-MP-41-01	ALKALINITY	W 125P	C4
X20031D-F	BE-MP15-01	METHANE	W 40V	2
X20032E-G	BE-MP17-01	METHANE	W 40V	2
X20033E-G	BE-MP-8-01	METHÀNE	W 40V	2
	BE-MP-41-01	METHANE	W 40V	2
	be returned GC X	Metals Wet Ch	nem X HPLC	SxPrep
	SxRec <u>C</u>	QA/QC <u>C</u> Acctg <u>C</u>	File <u>Ori</u> g	1
Page 1 of 1	L Page(s)		Cuntadian (nata)	MM 3/13/91

Custodian/Date: 1 1 3 1 3 196

CHAIN OF CUSTODY RECORD / ANALYTICAL SERVICES REQUEST

Page of	MICHAREL PHELPS	P.O.*	O' CI STD (2 wks) CI STD UST (3 day	U Other (Specify)	EAL use only Do not write	in shaded area	EAL	Project # 16 1/06	Custodian	EAL Sample N	Y	. 62.7	634	034		7	Location 2 C2/	Container Size
	3 PROJECT ID TE	1	TURNAHOUND REQUIRED*	C) Other (Speci	ANALYSIS REQUESTED	97/ 94/ 978M	NPDES / S	Metals	isis Metals Sircle & list Sircle & list Wood W		XXXX	メスススス	`	KKK K			12 5 26 12 3C	p d
Evergreen Analytical Inc.	4036 Younglield St. Wheat Ridge, Colorado 80033	(800) 425 0321 FAX (303) 425-6854 (800) 845-7400	FAX RESULTS Y / N		ANALYSIS RE	(circle)	508/508 (circle) 515 (circle) 715 (circle) 715 (circle) 715 (circle)	8080/6 8150/5 502 (cil	EPH 8015/ VPH 8015/ VPH 418-1/ VPH 418-1/ VPH 8050/ VPH 8050/ VPH 8015/ VPH									
	ST 200 💸	<	FAX # 510 769 9244	dille	×	SisieA	PesvHerb/A	\AN8\ S3\\$S8	WOOD THE	X	X	X	>			4	52	1
	AGE PEWY	10576 diz	1 FAX # 510	Wiles	MAŤRIX	puno.			Mo. of Conf Water-Duni (circle) Soil / Solid	X 9	30 7 X	15 PX	16'n. 3 K					
	MANA VIII	ATE CA_ZIP		Breeze	Krieden	S S	RINT	nation:	DATE SAMPLED TII	3/11/96 1040	3/11/16 230	3/11/6 15115	2/11/61	1				
	ADDRESS 1301 NACINA VILLAGE PLWY ST 200	CITY ALAINED A STATE	PHONE \$10 769 0100	Sampler Name: (signature) Profit of the signature) Profit of the signature	(print) Mystace H. Frieden	Evergreen Analytical Cooler N Cooler Received	Please PRIN	all information:	CLIENT SAMPLE IDENTIFICATION	BE. WP 15-\$1	BE-NP17-201	AT THE STATE OF TH	RK-M5-41-61			HT:	DD:	Instructions:







4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled : 3/7,8/96 **Date Received** : 3/9/96 **Date Prepared**

Date Analyzed

: 3/9/96

: 3/9/96

Client Project ID.

: 722450. 14020

Method

Lab Project Number: 96-0686 : EPA 300.0

Detection Limit

: 0.056 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Nitrate-N		Dilution <u>Factor</u>
X19943	BE-MP20-01	Water	2.6	mg/L	1
X19944	BE-MW02-01	Water	2.2	mg/L	1
X19945	BE-MP21-01	Water	1.8	mg/L	1
X19946	BE-MW01-01	Water	3.6	mg/L	1
X19947	BE-MW18-01	Water	2.3	mg/L	1
X19948	BE-MW19-01	Water	2.6	mg/L	1
Method Blank	(3/9/96)		<0.056	mg/L	

Quality Assurance*

	:	Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19943	BE-MP20-01 Matrix Spike	10.0	11.5	21.8	102
X19943	BE-MP20-01 Matrix Spike Du	10.0 p	11.5	21.8	102
MS/MSD F	RPD .				0.49

^{*} Quality Assurance results reported as Nitrate (NO₃)

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled : 3/7,8/96 Client Project ID. : 722450. 14020 **Date Received** : 3/9/96 Lab Project Number: 96-0686 Date Prepared : 3/9/96 Method : EPA 300.0 Date Analyzed : 3/9/96 **Detection Limit** : 0.25 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Sufate</u>		Dilution <u>Factor</u>
X19943	BE-MP20-01	Water	11.6	mg/L	1
X19944	BE-MW02-01	Water	8.4	mg/L	1
X19945	BE-MP21-01	Water	9.3	mg/L	1
X19946	BE-MW01-01	Water	26.8	mg/L	1
X19947	BE-MW18-01	Water	7.0	mg/L	1
X19948	BE-MW19-01	Water	25.9	mg/L	1
Method Blank	(3/9/96)		<0.25	mg/L	

Quality Assurance

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19943	BE-MP20-01 Matrix Spike	10.0	11.6	21.8	102
X19943	BE-MP20-01 Matrix Spike Du	10.0 IP	11.6	21.5	99
MS/MSD	RPD				0.03

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled : 3/7,8/96 **Date Received** : 3/9/96

Client Project ID. : 722450. 14020

Date Prepared : 3/9/96 Date Analyzed : 3/9/96 Lab Project Number: 96-0686 Method

: EPA 300.0

Detection	Limit
-----------	-------

: 0.25mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Chloride	Dilution <u>Factor</u>
X19943	BE-MP20-01	Water	25.1 mg	/L 1
X19944	BE-MW02-01	Water	13.5 mg	/L 1
X19945	BE-MP21-01	Water	23.3 mg/	/L 1
X19946	BE-MW01-01	Water	74.6 mg/	L 10
X19947	BE-MW18-01	Water	10.9 mg/	'L 1
X19948	BE-MW19-01	Water	103 mg/	L 10
Method Blank	(3/9/96)		<0.25 mg/	'L

Quality Assurance

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19943	BE-MP20-01 Matrix Spike	10.0	25.1	36.1	110
X19943	BE-MP20-01 latrix Spike Du	10.0 p	25.1	34.9	98
MS/MSD RPD					11.4

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

 Date Sampled
 : 3/7,8/96
 Client Project ID.
 : 722450. 14020

 Date Received
 : 3/9/96
 Lab Project Number
 : 96-0686

 Date Prepared
 : 3/9/96
 Method
 : EPA 300.0

 Date Analyzed
 : 3/9/96
 Detection Limit
 : 0.076 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Nitrite-N	Dilution <u>Factor</u>
X19943	BE-MP20-01	Water	<0.076 mg/L	1
X19944	BE-MW02-01	Water	<0.076 mg/L	1
X19945	BE-MP21-01	Water	<0.076 mg/L	1
X19946	BE-MW01-01	Water	<0.076 mg/L	1
X19947	BE-MW18-01	Water	<0.076 mg/L	1
X19948	BE-MW19-01	Water	<0.076 mg/L	1
Method Blank	(3/9/96)		<0.076 mg/L	

Quality Assurance*

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19943	BE-MP20-01 Matrix Spike	10.0	<0.25	10.0	100
X19943	BE-MP20-01 Matrix Spike Du	10.0 p	<0.25	10.0	100
MS/MSD R	RPD				0.10

^{*}Quality Assurance results reported as Nitrite (NO₂)

Analyst

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

Date Sampled : 3/7,8/96 **Date Received** : 3/9/96

Client Project ID. 96-0686

722450.14020

Date Prepared

: 3/11/96

Lab Project Number Method

EPA 310.1

Date Analyzed

: 3/11/96

Detection Limit

5.0 mgCaCO₃/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Total <u>Alkalinity</u> (mgCaCO ₃ /L)	Dilution <u>Factor</u>
X19943	BE-MP20-01	Water	84.6	1
X19944	BE-MW02-01	Water	86.4	1
X19945	BE-MP21-01	Water	76.8	1
X19946	BE-MW01-01	Water	177	1
X19947	BE-MW18-01	Water	67.6	1
X19948	BE-MW19-01	Water	91.9	1

Method Blank

(3/11/96)

< 5.0

Quality Assurance

Reference	<u>True Value</u> (mgCaCO ₃ /L)	Result (mgCaCO ₃ /L)	% Recovery
ERA Alkalinity Lot# 0814-95-0	120	116	97

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

Date Sampled

: 3/7,8/96

Client Project ID.

: 722450.14020

Date Received

: 3/9/96

Lab Project Number: 96-0686

Date Prepared

: 3/11/96

Method

: Standard Method 4500 NH₃ F

Date Analyzed

: 3/11/96

Detection Limit

: 0.80 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Ammonia-N	Dilution <u>Factor</u>
X19943	BE-MP20-01	Water	<0.80 mg/L	1
X19944	BE-MW02-01	Water	<0.80 mg/L	1
X19945	BE-MP21-01	Water	<0.80 mg/L	1
X19946	BE-MW01-01	Water	<0.80 mg/L	1
X19947	BE-MW18-01	Water	<0.80 mg/L	1
X19948	BE-MW19-01	Water	<0.80 mg/L	1

Method Blank

(3/11/96)

< 0.80 mg/L

Quality Assurance

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19943	BE-MP20-01 Matrix Spike	10.0	<0.80	9.1	91
X19943	BE-MP20-01 Matrix Spike Dup	10.0	<0.80	9.1	91

MS/MSD RPD

0.44

Evergreen Analytical, Inc. 4036 Youngfield, Wheat Ridge, CO 80033 (303) 425-6021

RSK-175 Gas Method Methane LCS Report Form

LCS No.

: LCS031396

EPA Method No.

: RSKSOP-175

Date Prepared

: 3/13/96

Matrix

: Water

Date Analyzed

: 3/13/96

Method Blank

: GB031396

E.A. LCS Source No.

: 1723

Lab File No.

: GAS0312038

	Spike	Method Blank	LCS		QC
Compound	Added	Concentration	Concentration	LCS	Limits
	(ug)	(ug)	(ug)	%REC	%REC
Methane Gas	500	0	361	72	67-85

Spike Recovery:

O out of (1) outside limits.

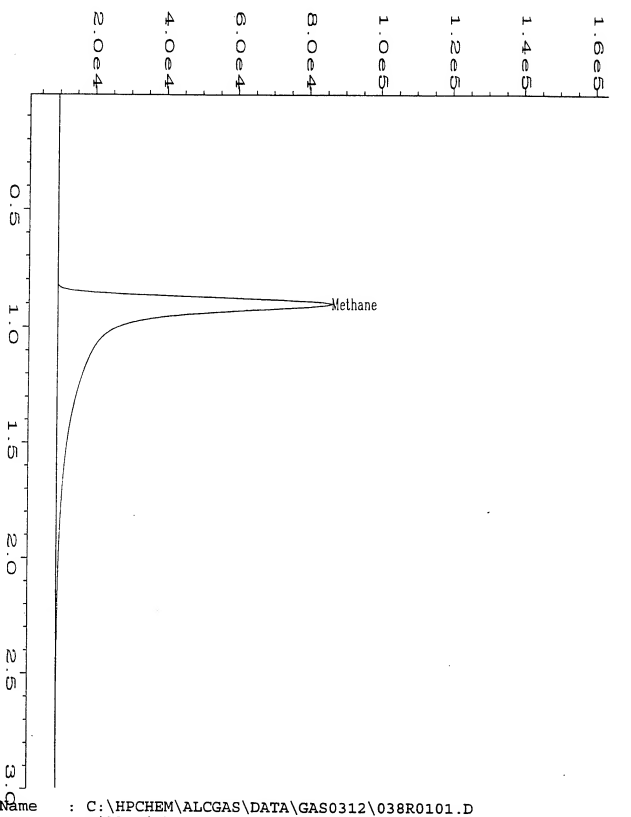
Note: The LCS was made by taking the sample and displacing 4ml of headspace with a 1%methane gas and shaking the VOA for 5 minutes. Then injecting 50 ul from the headspace into the GC resulting in a theoretical concentration of 500 ug.

NOTES:

* = Values outside of QC limits.

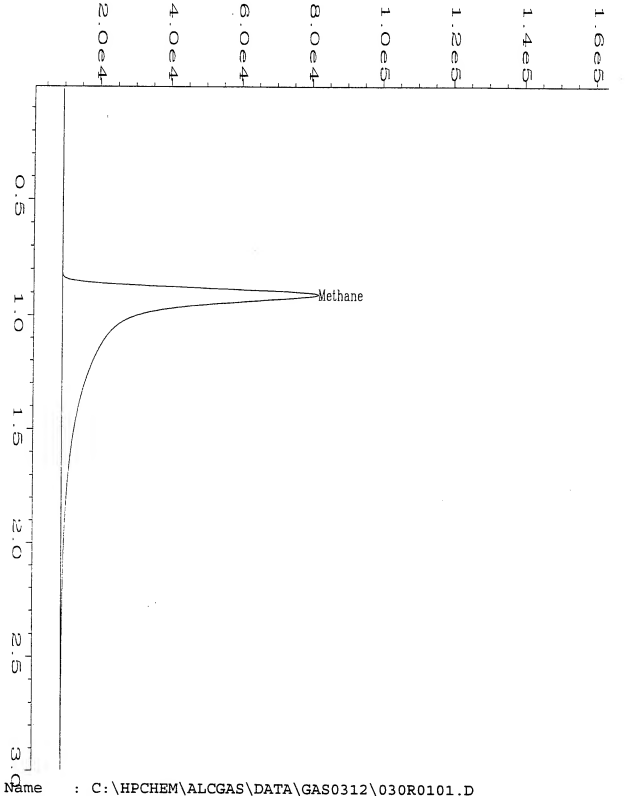
NA = Not analyzed/not available.

LCS0313.XLS; 3/13/96

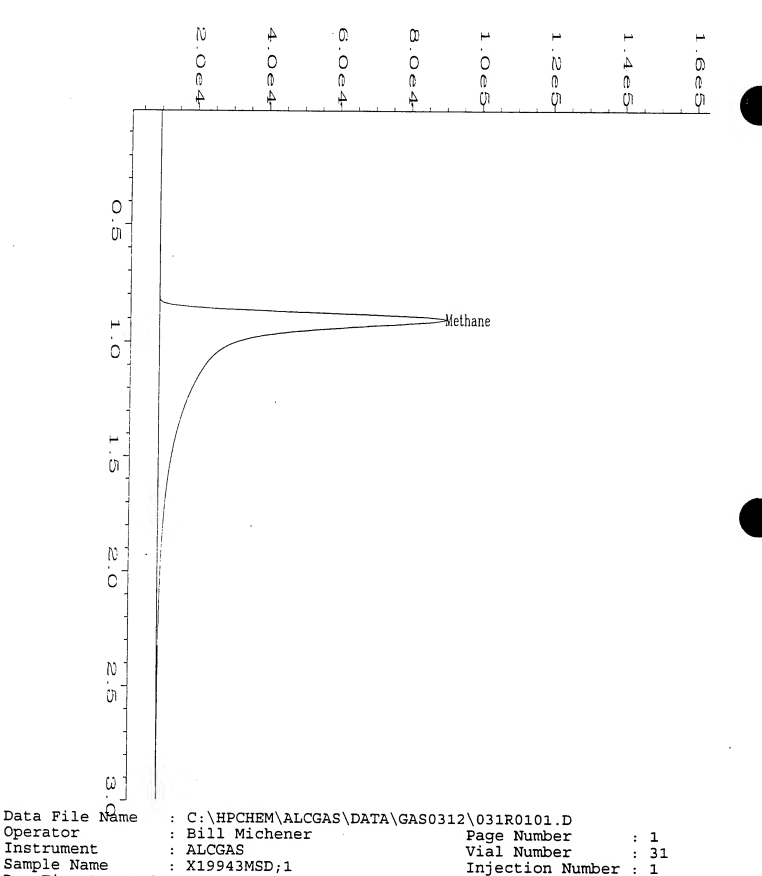


Data File Name :	C:\HPCHEM\ALCGAS\DATA\GAS0312	2\038R0101.D	
Operator :		Page Number	: 1
	ALCGAS		: 38
Sample Name :	LCS031396;Gas	Injection Number	
Run Time Bar Code:		Sequence Line	
	13 Mar 96 11:23 AM	Instrument Method:	: METHANE.MT
	13 Mar 96 11:26 AM	Analysis Method	: GAS0313.MT
Last Decalib on .	12 MAD OC OO. 22 AM	A	

Report Created on: 13 Mar 96 11:26 AM Analysis Method: M. Last Recalib on : 13 MAR 96 08:33 AM Sample Amount : 0 Multiplier : 1 ISTD Amount :



Data File Name :	C:\HPCHEM\ALCGAS\DATA\GAS031	2\030R0101.D
Operator :	Bill Michener	Page Number : 1
	ALCGAS	Vial Number : 30
Sample Name :		Injection Number : 1
Time Bar Code:		Sequence Line : 1
	13 Mar 96 09:38 AM	Instrument Method: METHANE.MTH
Rest Created on:	13 Mar 96 11:20 AM	Analysis Method : GAS0313.MTH
Recalib on :	13 MAR 96 08:33 AM	Sample Amount : 0
•	1	ISTD Amount :
Sample Info :	96-0686;BE-MP20-01;Displaced	4ml with 1% methane
	gas(#1723), shook for 5 min.	and injected 50ul to equal a
	theoretical spike of 500mg	The sample is injected at a



Run Time Bar Code: Sequence Line : 1 Acquired on : 13 Mar 96 09:45 AM Instrument Method: METHANE.M', Report Created on: 13 Mar 96 11:20 AM Analysis Method : GAS0313.M Last Recalib on : 13 MAR 96 08:33 AM Sample Amount Multiplier ISTD Amount Sample Info 96-0686; BE-MP20-01; Displaced 4ml with 1% methane gas(#1723), shook for $\bar{5}$ min. and injected 50ul to equal a theoretical spike of 500mg. The sample is injected at a

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Data File Name
                    : C:\HPCHEM\ALCGAS\DATA\GAS0312\029R0101.D
                    : Bill Michener
                                                        Page Number
Instrument
                    : ALCGAS
                                                        Vial Number
                                                                            : 29
Sample Name
                    : X19948Dup;1
                                                        Injection Number: 1
    Time Bar Code:
                                                        Sequence Line
   uired on
                    : 13 Mar 96
                                   09:27 AM
                                                        Instrument Method: METHANE.MTH
```

Analysis Method : GAS0313.MTH

: 0

Sample Amount

ISTD Amount

Multiplier 1 : 96-0686;BE-MW19-01;Water Sample Info

Recalib on : 13 MAR 96 08:33 AM

11:20 AM

t Created on: 13 Mar 96

Operator

Evergreen Analytical, Inc. 4036 Youngfield, Wheat Ridge, CO 80033 (303) 425-6021

RSK-175 Gas Method Methane, Ethane, Ethene Gas Matrix Spike / Matrix Spike Duplicate Report

Client Sample No.

: BE-MP20-01

Client Project No.

: 722450.14020

Lab Sample No.

: X19943

Lab Project No. EPA Method No.

: 96-0686 : RSKSOP-175

Date Sampled Date Received

: 3/8/96

Matrix

: Water

Date Prepared

: 3/9/96: 3/13/96

Method Blank

: GB031396

Date Analyzed

: 3/13/96

Lab File No's.

: GAS0312030,031

E.A. MS/MSD Spike Source No.

: 1723

	Spike	Sample	MS		QC
Compound	Added	Concentration	Concentration	MS	Limits
	(ug)	(ug)	(ug)	%REC	%REC
Methane Gas	500	0	391	78	40-89

Compound	Spike Added	MSD Concentration MSD		RPD	QC Limits	
	(ug)	(ug)	%REC		RPD	%REC
Methane Gas	500	428	86	9.0	0-24.4	40-89

Spike Recovery:

0 out of (1) outside limits.
0 out of (2) outside limits.

NOTES:

* = Values outside of QC limits.

NA = Not analyzed/not available

Note: The Spike was made by taking the sample and displacing 4ml of headspace with a 1% methane gas and shaking the VOA for 5 minutes. Then injecting 50 ul from the headspace into the GC resulting in a theoretical concentration of 500 ug.

Analyst

Approved

MS0686.XLS; 3/19/96

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                      : C:\HPCHEM\ALCGAS\DATA\GAS0312\028R0101.D
                      : Bill Michener
                                                              Page Number
                                                                                     : 1
```

Operator Instrument : ALCGAS Vial Number 28 Sample Name : X19948;1 Injection Number: 1 Time Bar Code: Sequence Line _uired on : 13 Mar 96 09:23 AM Instrument Method: METHANE.MTH Created on: 13 Mar 96 Re 11:20 AM Analysis Method : GAS0313.MTH lecalib on : 13 MAR 96 08:33 AM Sample Amount : 0 Multiplier 1 ISTD Amount :

Sample Info : 96-0686;BE-MW19-01;Water

Methane Report Form

Client Sample Number Lab Sample Number Date Sampled Date Received Date Extracted/Prepared Date Analyzed	: BE-MW19-01 : X19948Dup : 3/7/96 : 3/9/96 : 3/13/96	Client Project No. Lab Project No. Dilution Factor Method Matrix Lab File No.	: 722450.14020 : 96-0686 : 1.00 : RSKSOP-175 : Water : GAS0312029
Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002
		•	
Temperature Amount Injected	: <u>69.6</u> : 0.5		Meth
Total Volume of Sample Head space created Methane Area	: 43 : 4 : 0	ml Concentration ml in Head Space	Meth

16 g

QUALIFIERS:

E = Extrapolated value.

Atomic weight(Methane)

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

K. Hollman

Analyst

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Data File Name
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                     : Bill Michener
                                                            Page Number
                                                                                  : 1
                     : ALCGAS
                                                            Vial Number
                                                                                  : 27
                     : X19947;1
                                                            Injection Number: 1
```

Operator Instrument Sample Name Time Bar Code: Sequence Line : 1 uired on : 13 Mar 96 09:19 AM Instrument Method: METHANE.MTH Created on: 13 Mar 96 Analysis Method 11:20 AM : GAS0313.MTH ecalib on : 13 MAR 96 08:33 AM Sample Amount : 0 Multiplier : 1 ISTD Amount Sample Info : 96-0686;BE-MW18-01;Water

Methane Report Form

mg/L

U

Compound Name	Cas Number	Sample Concentration	RL
Date Analyzed	: 3/13/96	Lab File No.	: GAS0312028
Date Extracted/Prepared	: 3/13/96	Matrix	: Water
Date Received	: 3/9/96	Method	: RSKSOP-175
Date Sampled	: 3/7/96	Dilution Factor	: 1.00
Lab Sample Number	: X19948	Lab Project No.	: 96-0686
Client Sample Number	: BE-MW19-01	Client Project No.	: 722450.14020

74-82-8

Temperature Amount Injected Total Volume of Sample Head space created	:	69.7 F 0.5 ml 43 ml 4 ml	Saturation Concentration Concentration in Head Space	Meth Meth	
Methane Area	:	O ug			

Atomic weight(Methane) 16 g

QUALIFIERS:

Methane

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

mg/L

0.002

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Instrument
                     : ALCGAS
                                                          Vial Number
Sample Name
                     : X19946;1
                                                          Injection Number :
                                                                                 1
    Time Bar Code:
                                                          Sequence Line
                                                                                 1
   nuired on
                    : 13 Mar 96
                                    09:16 AM
                                                          Instrument Method: METHANE.MTH
```

Analysis Method

Sample Amount

ISTD Amount

: GAS0313.MTH

: 0

Multiplier Sample Info

Operator

: 1 : 96-0686; BE-MW01-01; Water

: 13 MAR 96 08:33 AM

11:20 AM

Created on: 13 Mar 96

Recalib on

Methane Report Form

Client Sample Number Lab Sample Number Date Sampled Date Received Date Extracted/Prepared Date Analyzed	: BE-MW18-01 : X19947 : 3/7/96 : 3/9/96 : 3/13/96 : 3/13/96	Client Project No. Lab Project No. Dilution Factor Method Matrix Lab File No.	: 722450.14020 : 96-0686 : 1.00 : RSKSOP-175 : Water : GAS0312027
Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002
Temperature Amount Injected	: 66.7 I	nl Concentration	eth
Total Volume of Sample Head space created Methane Area	: 43 r : 4 r	nl <u>in Head Space</u>	eth

16 g

QUALIFIERS:

E = Extrapolated value.

Atomic weight(Methane)

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

K. Hollman

AF0686.XLS

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Operator Instrument : ALCGAS Vial Number 25 Sample Name : X19945;1 Injection Number Time Bar Code: Sequence Line : 1 <u>uired</u> on : 13 Mar 96 09:12 AM Instrument Method: METHANE.MTH Created on: 13 Mar 96 11:20 AM Analysis Method : GAS0313.MTH Recalib on : 13 MAR 96 08:33 AM Sample Amount ISTD Amount

Multiplier : 1 : 96-0686;BE-MP21-01;Water Sample Info

Methane Report Form

Client Sample Number Lab Sample Number Date Sampled Date Received	: BE-MW01-01	Client Project No.	: 722450.14020
	: X19946	Lab Project No.	: 96-0686
	: 3/8/96	Dilution Factor	: 1.00
	: 3/9/96	Method	: RSKSOP-175
Date Extracted/Prepared Date Analyzed	: 3/13/96 : 3/13/96	Matrix Lab File No.	: RSKSOP-175 : Water : GAS0312026

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

Temperature Amount Injected Total Volume of Sample Head space created		66.6 F 0.5 ml 43 ml 4 ml	Saturation Concentration Concentration in Head Space	Meth Meth		
Methane Area	:	0 ug			**************************************	

Atomic weight(Methane) : _____ <u>16</u> g

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

K. Hollman
Approved

Analyst

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Instrument
                   : ALCGAS
                                                      Vial Number
                                                                          : 24.
Sample Name
                   : X19944;1
                                                      Injection Number : 1
    Time Bar Code:
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                                                                          : 1
   lired on
                   : 13 Mar 96
                                                      Instrument Method: METHANE.MTH
                                  09:08 AM
Report Created on: 13 Mar 96
                                  11:20 AM
                                                      Analysis Method : GAS0313.MTH
       ecalib on
                  : 13 MAR 96 08:33 AM
                                                      Sample Amount
                                                                         : 0
      plier
                   : 1
                                                      ISTD Amount
                   : 96-0686;BE-MW02-01;Water
Sample Info
```

Operator

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Mul

Methane Report Form

Client Sample Number	: BE-MP21-01	Client Project No.	: 722450.14020
Lab Sample Number	: X19945	Lab Project No.	: 96-0686
Date Sampled	: 3/8/96	Dilution Factor	: 1.00
Date Received	: 3/9/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/13/96	Matrix	: Water
Date Analyzed	: 3/13/96	Lab File No.	: GAS0312025

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

Temperature Amount Injected Total Volume of Sample Head space created		65.3 F 0.5 ml 43 ml 4 ml	Saturation Concentration Concentration in Head Space	Meth Meth	
Methane Area	•	0 ug			

Atomic weight(Methane) : ______ 16 g

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

K Hollman
Approved

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Operator :	Bill Michener	Page Number :	1
Instrument :		Vial Number :	
Sample Name :	X19943;1	Injection Number :	
F Time Bar Code:		Sequence Line :	
_uired on :	13 Mar 96 09:04 AM	Instrument Method:	
keret Created on:	13 Mar 96 11:20 AM	Analysis Method :	
Recalib on :	13 MAR 96 08:33 AM	Sample Amount :	0
	1	ISTD Amount :	
Sample Info :	96-0686;BE-MP20-01;Water	•	

Methane Report Form

Client Sample Number Lab Sample Number Date Sampled Date Received Date Extracted/Prepared Date Analyzed	: BE-MW02-01 : X19944 : 3/8/96 : 3/9/96 : 3/13/96	Client Project No. Lab Project No. Dilution Factor Method Matrix Lab File No.	: 722450.14020 : 96-0686 : 1.00 : RSKSOP-175 : Water : GAS0312024
Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

67.4 F

0.5 ml

43 ml

4 m!

0 ug

Saturation

Concentration

Concentration

in Head Space

Meth

Meth

Atomic weight(Methane)	:	16 g

QUALIFIERS:

Temperature

Amount Injected

Methane Area

Total Volume of Sample

Head space created

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

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                     : ALCGAS
                                                          Vial Number
                                                                               : 22
                     : GB031396
                                                          Injection Number : 1
```

Operator Instrument Sample Name Time Bar Code: Sequence Line : 1 red on : 13 Mar 96 08:45 AM Instrument Method: METHANE.MTH Re Created on: 13 Mar 96 11:20 AM Analysis Method : GAS0313.MTH Recalib on : 13 MAR 96 08:33 AM Sample Amount : 0 Multiplier : 1 ISTD Amount Sample Info : Gas Method Blank

Displaced 4ml of deionized water in 43ml vial with Helium, shook for 5 min and injected 500ml

Methane Report Form

Client Sample Number Lab Sample Number Date Sampled Date Received Date Extracted/Prepared Date Analyzed	: BE-MP20-01 : X19943 : 3/8/96 : 3/9/96 : 3/13/96 : 3/13/96	Client Project No. Lab Project No. Dilution Factor Method Matrix Lab File No.	: 722450.14020 : 96-0686 : 1.00 : RSKSOP-175 : Water : GAS0312023
Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002
Temperature	:66.5		Meth
Amount Injected	:0.5		
Total Volume of Sample	: 43		Meth
Head space created		ml <u>in Head Space</u>	
Methane Area	:0	ug	

<u>16</u> g

QUALIFIERS:

E = Extrapolated value.

Atomic weight(Methane)

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

K Hollman

Analyst

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

Date Sampled : 3/7,8/96 **Date Received** : 3/9/96

Client Project ID. Lab Project Number: 96-0686

: 722450. 14020

0.55

Date Prepared

Method

: EPA 415.1

Date Analyzed

: 3/13/96 : 3/13/96

Detection Limit

: 1.0 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Total Organic <u>Carbon</u>	Dilution <u>Factor</u>
X19943	BE-MP20-01	Water	1.2 mg/L	1
X19944	BE-MW02-01	Water	1.6 mg/L	1
X19945	BE-MP21-01	Water	1.8 mg/L	1
X19946	BE-MW01-01	Water	7.3 mg/L	1
X19947	BE-MW18-01	Water	2.2 mg/L	1
X19948	BE-MW19-01	Water	3.2 mg/L	1

Method Blank

(3/13/96)

<1.0

Quality Assurance

	4	Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19943	BE-MP20-01 Matrix Spike	10.0	1.2	12.1	109
X19943	BE-MP20-01 Matrix Spike Du	10.0	1.2	12.1	109
MS/MSD R	PD				0.55

Approved

Methane Report Form Method Blank Report

Method Blank Number

: GB031396

Client Project No.

: 722450.14020

Date Extracted/Prepared

: 3/13/96

Lab Project No.

: 96-0686

Date Analyzed

: 3/13/96

Dilution Factor

: 1.00

Method

Matrix

: RSKSOP-175

: Water

Lab File No.

: GAS0312022

_		Sample	
Compound Name	Cas Number	Concentration	RL
		mg/L	mg/L
Methane	74-82-8	U	0.002

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Page / of **IALYTICAL SERVICES REQUEST** en Analytical Inc. CHAIN OF CUSTODY RECORD Ever 11.4° # 200 MARSONS ES

4036 Younglield St. Wheat Ridge, Colorado 80033 (303) 425-6021 FAX (303) 425-6854 (800) 845-7400 FAX RESULTS Y / N FAX#510 769 9244 PHONE # 510769 0100 ADDRESS 130 MARCINA CITY PLANKED A STATE CA COMPANY

PATTY LICCIELLAN	150,14020	P.O.#
CLIENT CONTACT (print)_	PROJECT I.D. 72245	EAL. QUOTE #

TURNAROUND REQUIRED* 🗆 STD (2 wks) 🗀 STD UST (3 day)

Sampler Name: . (signature) That Manager					*ex	 Other (Speciexpedited turnaround subject to additional fee 	Oth	Other (Specify)
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Evergreen Analytical Cooler NoCooler Received	icound		أدداه)			Darlo		in shaded area
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all information:	<u>α</u> ⁄δυιγ	950	8080/ 8150/	Oli & C	DOM: Powers	201		Project # / O O O O
CLIENT SAMPLE DATE IDENTIFICATION SAMPLED TIME 2	Mo. of Con Water-Drin (circle) Soil / Solid Oil / Sludge	TCLP VOA (circle) VOA 8260/	Pesticides Pest/PCBs Herbicides PCB Scree	0208 X3T8 7.814 H9AT 7.19PH 8015	TEPH 8015 Total Metali Circle & list Circle & list Circle & list	MW III		
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BE-NP21-01 3/8/96 1320-	1 X H	X			7	X X X X		45
BE- MWD1-01 3/8/96 1430 7	717	メ			X	XXX		46
SE-MW18-01/3/7/96/15007	4 X	メ			\ <u>\</u>	X X X		47
BE-MW19-01 3/7/26 1345 6	X	X			×	XX		48
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DO:							ŏ	Container Size (25.4 40.7)
Instructions: Analyze for PE-MW19-01		alkalinity a	and ammonia	يد ا	Sufficient	John Hombs) amolow	7
1 96 n		-			X	ł		

Date/Time Relinquished by: (Signature)

Date/Time Received by: (Signature)

Date/Time

Evergreen Analytical Sample Receipt/Check-in Record Date & Time Rec'd: 3/9/46 1105 Shipped Via: Fedex Client : Jaruns Client Project ID(s): 722450_/4020 EAL Project #(s):96- 0686 EAL Cooler(s): Y NA Cooler# Ø Ice packs N Y N Y N Y N Y N Temperature ℃ _ 6 > 8 N N/A 1. Custody seal(s) present: Seals on cooler intact Seals on bottle intact 2. Chain of Custody present: 3. Samples Radioactive: (Comment on COC if >0.5mr/h) 4. Containers broken or leaking: (Comment on COC if Y) 5. Containers labeled: 6. COC agrees w/ bottles received: (Comment on COC if N) 7. COC agrees w/ labels: (Comment on COC if N) 8. Headspace in vials-waters only: (Comment on COC If 1) 9. VOA samples preserved: 10. pH measured on metals, cyanide or phenolics*: List discrepancies_ *Non-EAL provided containers only, water samples only. 11. Metal samples present: Total _____, Dissolved _____, TCLP ____ D or PD to be filtered: T,TR,D,PD to be Preserved: 12. Short holding times: Specify parameters_____ 13. Multi-phase sample(s) present: 14. COC signed w/ date/time: Comments: Cooper belongs to ITS ANAMETRIX, 1961 Concum-sx De Sauture CA 95/31

(Additional comments on back)

Custodian Signature/Date:_

JA 3/4/96

Evergreen Analytical Sample Log Sheet	Project # <u>96-0686</u>			
Date(s) Sampled: 03/07,08/96 COC	Date Due: 03/25/96			
Pate Received: 03/09/96 1105	Holding Time(s): 03/09,10-NO ₂ ,NO ₃ ,			
ent Project I.D. 722450.14020	3/21,22-ALKALINITY Rush STANDARD			
Client: PARSONS ENGINEERING SCIENCE, INC	C. Cooler Return 5.00			
Address: 1301 MARINA VILLAGE PKWY #200				
ALAMEDA, CA 94501	Airbill # FEDEX 1154092752			
Contact: MICHAEL FREIDMAN	Client P.O.			
Phone #510-769-0100 Fax #510-769-9244				
Special Invoicing/Billing				
Special Instructions				
Lab Client				
Lab Client ID# ID# Analysi	s Mtx Btl Loc			
X19943A BE-MP20-01 TOC	W 125A C4			
X19944A BE-MW02-01 TOC	W 125A C4			
X19945A BE-MP21-01 TOC	W 125A C4			
X19946A BE-MW01-01 TOC	W 125A C4			
947A BE-MW18-01 TOC	W 125A C4			
X10048A BE-MW19-01 TOC, AMMO	NIA W 125A C4			
X 3B-D BE-MP20-01 METHANE	W 40V C4			
X19944B-D BE-MW02-01 METHANE	W 40V C4			
X19945B-D BE-MP21-01 METHANE	W 40V C4			
X19946B-D BE-MW01-01 METHANE	W 40V C4			
X19947B-D BE-MW18-01 METHANE	W 40V C4			
X19948B-D BE-MW19-01 METHANE	W 40V C4			
X19943G BE-MP20-01 NO ₂ , NO ₃ , SO	4,Cl W 125P C4			
$X19944G \qquad BE-MW02-01 \qquad NO_2, NO_3, SO$	4,Cl W 125P C4			
<u>X19945G BE-MP21-01 NO₂, NO₃, SO</u>	4,Cl W 125P C4			
<u>X19946G BE-MW01-01 NO₂, NO₃, SO</u>	4,Cl W 125P C4			
<u>X19947G BE-MW18-01 NO₂, NO₃, SO</u>	4,C1 W 125P C4			
X19948G BE-MW19-01 NO ₂ , NO ₃ , SO ₄ , C	Cl ⁻ ,ALK. W 125P C4			
X19943E BE-MP20-01 AMMONIA	W 500P C4			
X19944E BE-MW02-01 AMMONIA	W 500P C4			
R=Sample to be returned te GC/MS GC X Metals	Wet Chem X HPLC _ SxPrep _			
SxRec \underline{C} QA/QC \underline{C} Ac	cctg <u>C</u> File <u>Orig</u>			
Page 1 of 2 Page(s)	Custodian/Date: 18 3/11/96 Pm 3/12/96			

Lab ID #	Client ID#	Analysis	Mtx	Btl Loc	
X19945E	BE-MP21-01	AMMONIA	W	500P C4	
X19946E	BE-MW01-01	AMMONIA	W	500P C4	
X19947E	BE-MW18-01	AMMONIA	W	500P C4	
X19943F	BE-MP20-01	ALKALINITY	W	125P C4	
X19944F	BE-MW02-01	ALKALINITY	W	125P C4	
X19945F	BE-MP21-01	ALKALINITY	W	125P C4	
X19946F	BE-MW01-01	ALKALINITY	W	125P C4	
X19947F	BE-MW18-01	ALKALINITY	W	125P C4	

Page 2 of 2 Pages Project <u># 96-0686</u>

R=Sample to be returned

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled

: 3/06-07/96

Client Project ID.

: 722450.14020 Beale

Date Received

: 3/08/96

Lab Project Number: 96-0673

Date Prepared

: 3/08/96

Method

: EPA 300.0

Date Analyzed

: 3/08/96

Detection Limit

: 0.25 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Sulfate</u>		Dilution <u>Factor</u>
X19900	BE-MP-27-01	Water	5.2	mg/L	1
X19901	BE-MP-14-01	Water	25.2	mg/L	1
X19902	BE-MP-26-01	Water	4.5	mg/L	1
X19903	BE-MP-24-01	Water	9.1	mg/L	1
X19904	BE-MP-03-01	Water	17.1	mg/L	1
X19905	BE-MP-25-01	Water	5.8	mg/L	1
Method Blank	(3/8/96)	Water	< 0.25	ma/L	1

Quality Assurance

	<u>Sr</u>	nike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19900	Matrix Spike	10.0	5.2	13.8	86
X19900	Matrix Spike Dup	10.0	5.2	14.0	88
MS/MSD	RPD				1.8

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled

: 3/06-07/96

Client Project ID.

: 722450.14020 Beale

Date Received

: 3/08/96

Lab Project Number: 96-0673

50-0073 504 000 0

Date Prepared

: 3/08/96

Method

: EPA 300.0 -

Date Analyzed

: 3/08/96

Detection Limit

: 0.076 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	<u>Nitrite-N</u>	Dilution <u>Factor</u>
X19900	BE-MP-27-01	Water	<0.076 mg/L	1
X19901	BE-MP-14-01	Water	<0.076 mg/L	1 1
X19902	BE-MP-26-01	Water	<0.076 mg/L	1
X19903	BE-MP-24-01	Water	<0.076 mg/L	. 1
X19904	BE-MP-03-01	Water	<0.076 mg/L	1
X19905	BE-MP-25-01	Water	<0.076 mg/L	1
Method Blank	(3/8/96)	Water	<0.076 mg/L	1

Quality Assurance

	<u>Sr</u>	nike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19900	Matrix Spike	10.0	<0.25	10.0	100
X19900	Matrix Spike Dup	10.0	<0.25	9.6	96
MS/MSD F	RPD				4.2

Analyst Analyst

Approve

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled

: 3/06-07/96

Client Project ID.

: 722450.14020 Beale

Date Received Date Prepared

: 3/08/96

Lab Project Number: 96-0673

: EPA 300.0

: 3/08/96

Method

Date Analyzed

: 3/08/96

Detection Limit

: 0.056 mg/L

F	0 11				
Evergreen Sample #	Client <u>Sample ID.</u>	<u>Matrix</u>	Nitrate-N		Dilution <u>Factor</u>
X19900	BE-MP-27-01	Water	1.8	mg/L	1
X19901	BE-MP-14-01	Water	4.3	mg/L	1
X19902	BE-MP-26-01	Water	1.6	mg/L	1
X19903	BE-MP-24-01	Water	2.6	mg/L	1
X19904	BE-MP-03-01	Water	2.6	mg/L	1
X19905	BE-MP-25-01	Water	1.8	mg/L	1
Method Blank	(3/8/96)	Water	< 0.056	mg/L	1

Quality Assurance

	Si	oike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19900	Matrix Spike	10.0	8.1	17.5	94
X19900	Matrix Spike Dup	10.0	8.1	17.2	91
MS/MSD	RPD				3.0

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

Date Sampled : 3/6,7/96
Date Received : 3/8/96
Date Prepared : 3/11/96
Date Analyzed : 3/11/96

Client Project ID. : 722450.14020 Lab Project Number : 96-0673

Method : EPA 310.1

Detection Limit : 5.0 mg CaCO₃/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Total <u>Alkalinity</u> (mg CaCO ₃ /L)	Dilution <u>Factor</u>
X19900	BE-MP-27-01	Water	59.4	1
X19901	BE-MP-14-01	Water	287	1
X19902	BE-MP-26-01	Water	52.6	1
X19903	BE-MP-24-01	Water	52.8	1
X19904	BE-MP-03-01	Water	84.4	1
X19905	BE-MP-25-01	Water	60.4	1
Method Blank	(3/11/96)	Water	<5.0	1

Quality Assurance

Reference	<u>True Value</u> (mgCaCO ₃ /L)	$\frac{Result}{(mgCaCO_3/L)}$	% Recovery
ERA-Alkalinity	120	116	97

Analyst

Approved

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Anion Report

Date Sampled : 3/06-07/96 Date Received : 3/08/96

3/08/96

Client Project ID. : 722450.1 Lab Project Number : 96-0673

: 722450.14020 Beale

Date Received
Date Prepared

: 3/08/96

Method

: EPA 300.0

Date Analyzed

: 3/08/96

Detection Limit

: 0.25 mg/L

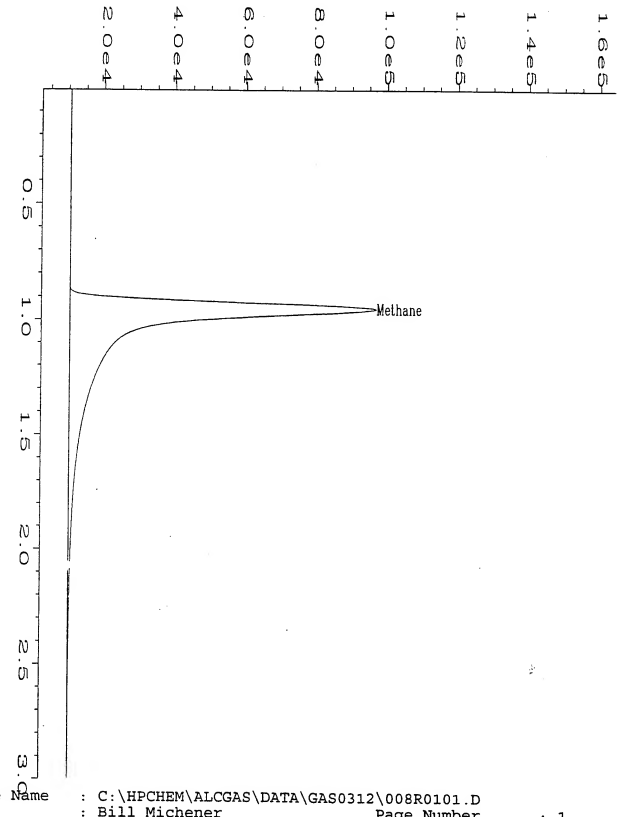
Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Chloride		Dilution <u>Factor</u>
X19900	BE-MP-27-01	Water	10.2	mg/L	1
X19901	BE-MP-14-01	Water	142	mg/L	10
X19902	BE-MP-26-01	Water	8.1	mg/L	1
X19903	BE-MP-24-01	Water	11.2	mg/L	1
X19904	BE-MP-03-01	Water	25.7	mg/L	1
X19905	BE-MP-25-01	Water	10.7	mg/L	1
Method Blank	(3/8/96)	Water	< 0.25	mg/L	1

Quality Assurance

	2	Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19900	Matrix Spike	10.0	10.2	19.2	91
X19900	Matrix Spike Dur	10.0	10.2	19.5	93
MS/MSD	RPD				2.6

Analyst

Approved



Data File Name Operator : Bill Michener Page Number Instrument : ALCGAS Vial Number Sample Name : LCS031296;Gas Injection Number : 1 1 Time Bar Code: Sequence Line : 1 Acquired on : 12 Mar 96 09:34 AM Instrument Method: METHANE.MTH Re Created on: 12 Mar 96 09:41 AM Analysis Method : GAS0312.MTH Recalib on : 12 MAR 96 09:33 AM Sample Amount : 0 Multiplier : 1 ISTD Amount Sample Info : Laboratory Control Sample

Displaced 4ml of deionized water in 43ml vial with 1% methane das(#1723) shook for 5 min and injected 50ml

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Analysis Report

: 722450.14020

Date Sampled : 3/6,7/96 Client Project ID.

Date Received : 3/8/96 Lab Project Numbe

Date Received : 3/8/96 Lab Project Number : 96-0673

Date Prepared : 3/8/96 Method : Standard Method 4500-NH₃F

Date Analyzed : 3/8/96 Detection Limit : 0.80 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Ammonia-N	Dilution <u>Factor</u>
X19900	BE-MP-27-01	Water	<0.80 mg/L	1
X19901	BE-MP-14-01	Water	<0.80 mg/L	1
X19902	BE-MP-26-01	Water	<0.80 mg/L	1
X19903	BE-MP-24-01	Water	<0.80 mg/L	1
X19904	BE-MP-03-01	Water	<0.80 mg/L	1
X19905	BE-MP-25-01	Water	<0.80 mg/L	1
Method Blank	(3/8/96)	Water	<0.80 mg/L	1

Quality Assurance

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19905	BE-MP-25-01 Matrix Spike	10.0	<0.80	10.0	100
X19905	BE-MP-25-01 Matrix Spike Du	p 10.0 .	<0.80	10.0	101
MS/MSD F	RPD				0.79

Analyst

Approved

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Instrument
                    : ALCGAS
                                                         Vial Number
                                                                              : 16
Sample Name
                      X19905;1
                                                         Injection Number: 1
   ` Time Bar Code:
                                                         Sequence Line
                                                                              : 1
   uired on
                    : 12 Mar 96
                                    11:52 AM
                                                         Instrument Method: METHANE.MTH
     t Created on: 13 Mar 96
                                    11:19 AM
                                                         Analysis Method
                                                                            : GAS0312.MTH
```

Sample Amount

ISTD Amount

Murciplier Sample Info : 96-0673; BE-MP-25-01; Water

: 12 MAR 96 09:33 AM

Operator

Recalib on

Evergreen Analytical, Inc. 4036 Youngfield, Wheat Ridge, CO 80033 (303) 425-6021

RSK-175 Gas Method Methane LCS Report Form

LCS No.

: LCS031296

EPA Method No.

: RSKSOP-175

Date Prepared

: 3/12/96

Matrix

: Water

Date Analyzed

: 3/12/96

Method Blank

: GB031296

E.A. LCS Source No.

: 1723

Lab File No.

: GAS0312008

	Spike	Method Blank	LCS		QC
Compound	Added	Concentration	Concentration	LCS	Limits
	(ug)	(ug)	(ug)	%REC	%REC
Methane Gas	500	0	466	93 *	67-85

Spike Recovery:

1 out of (1) outside limits.

Note: The LCS was made by taking the sample and displacing 4ml of headspace with a 1%methane gas and shaking the VOA for 5 minutes. Then injecting 50 ul from the headspace into the GC resulting in a theoretical concentration of 500 ug.

NOTES:

* = Values outside of QC limits.

NA = Not analyzed/not available.

LCS0312.XLS; 3/13/96

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Sample Name
                    : X19904;1
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   uired on
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                                                       Instrument Method: METHANE.MTH
    rt Created on: 13 Mar 96
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                                                       Analysis Method : GAS0312.MTH
     Recalib on
                  : 12 MAR 96 09:33 AM
                                                       Sample Amount
Mu. iplier
                                                       ISTD Amount
```

: 96-0673;BE-MP-03-01;Water

Operator

Sample Info

Re

Methane Report Form

Client Sample Number Lab Sample Number Date Sampled Date Received Date Extracted/Prepared Date Analyzed	: BE-MP-25-01 : X19905 : 3/7/96 : 3/8/96 : 3/12/96 : 3/12/96	Client Project No. Lab Project No. Dilution Factor Method Matrix Lab File No.	: 722450.14020 Beat : 96-0673 : 1.00 : RSKSOP-175 : Water : GAS0312016	
Compound Name	Cas Number	Sample Concentration mg/L	RL . mg/L	
Methane	74-82-8	U	0.002	
Temperature		2.9 F Saturation	Meth	`.
Amount Injected Total Volume of Sample Head space created Methane Area		0.5 mlConcentration43 mlConcentration4 mlin Head Space0 ug	Meth	

16 g

QUALIFIERS:

E = Extrapolated value.

Atomic weight(Methane)

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

K. Hollman

Analyst

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Sample Name
                    : X19903Dup;1
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                    : 12 Mar 96
                                   11:43 AM
                                                        Instrument Method: METHANE.MTH
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                                                        Analysis Method
                                                                           : GAS0312.MTH
                  : 12 MAR 96 09:33 AM
     Recalib on
                                                        Sample Amount
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Musciplier
```

ISTD Amount

Operator

Sample Info

: 1

: 96-0673;BE-MP-24-01;Water

EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP-03-01	Client Project No.	: 722450.14020 Bea
Lab Sample Number	: X19904	Lab Project No.	: 96-0673
Date Sampled	: 3/7/96	Dilution Factor	: 1.00
Date Received	: 3/8/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/12/96	Matrix	: Water
Date Analyzed	: 3/12/96	Lab File No.	: GAS0312015

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

Temperature	:	72.7 F	Saturation	Meth	
Amount Injected	;	0.5 ml	Concentration		-
Total Volume of Sample	:	43 ml	Concentration	Meth	
Head space created	.;	4 mi	in Head Space		
Methane Area	:	0 ug			

Atomic weight(Methane) : _____ g

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

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Approved

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         : ALCGAS
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Data File Name Operator Instrument Vial Number : 13 Sample Name : X19903;1 Injection Number: 1 Time Bar Code: Sequence Line duired on : 12 Mar 96 Instrument Method: METHANE.MTH rt Created on: 13 Mar 96 11:19 AM Analysis Method : GAS0312.MTH

Recalib on : 12 MAR 96 09:33 AM Sample Amount : 0

Murtiplier : 1 ISTD Amount :

Sample Info : 96-0673;BE-MP-24-01;Water

EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP-24-01	Client Project No.	: 722450.14020 Beal
Lab Sample Number	: X19903Dup	Lab Project No.	: 96-0673
Date Sampled	: 3/6/96	Dilution Factor	: 1.00
Date Received	: 3/8/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/12/96	Matrix	: Water
Date Analyzed	: 3/12/96	Lab File No.	: GAS0312014

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

Temperature	:	72.8 F	Saturation	Meth	
Amount Injected	•	0.5 ml	Concentration		
Total Volume of Sample	:	43 ml	Concentration	Meth	
Head space created	:	4 ml	in Head Space		
Methane Area	:	O uo			

Atomic weight(Methane)	:	16 g

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Analyst

Hollman Approved

AF0673.XLS

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Instrument
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Sample Name
                   : X19902;1
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                   : 12 Mar 96
                                 11:18 AM
                                                      Instrument Method: METHANE.MTH
    rt Created on: 13 Mar 96
                                 11:19 AM
                                                      Analysis Method
                                                                         : GAS0312.MTH
```

Muliplier Sample Info : 96-0673;BE-MP-26-01;Water

: 12 MAR 96 09:33 AM

Operator

Recalib on

Re

Sample Amount : 0 ISTD Amount

EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP-24-01	Client Project No.	: 722450.14020 B
Lab Sample Number	: X19903	Lab Project No.	: 96-0673
Date Sampled	: 3/6/96	Dilution Factor	: 1.00
Date Received	: 3/8/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/12/96	Matrix	: Water
Date Analyzed	: 3/12/96	Lab File No.	: GAS0312013

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

Temperature	:	72.6 F	Saturation	Meth	
Amount Injected	: <u></u>	0.5 ml	Concentration		-
Total Volume of Sample	:	43 ml	Concentration	Meth	
Head space created	:	4 ml	in Head Space		
Methane Area	:	0 ug			

16 g

QUALIFIERS:

E = Extrapolated value.

Atomic weight(Methane)

U = Compound analyzed for, but not detected.

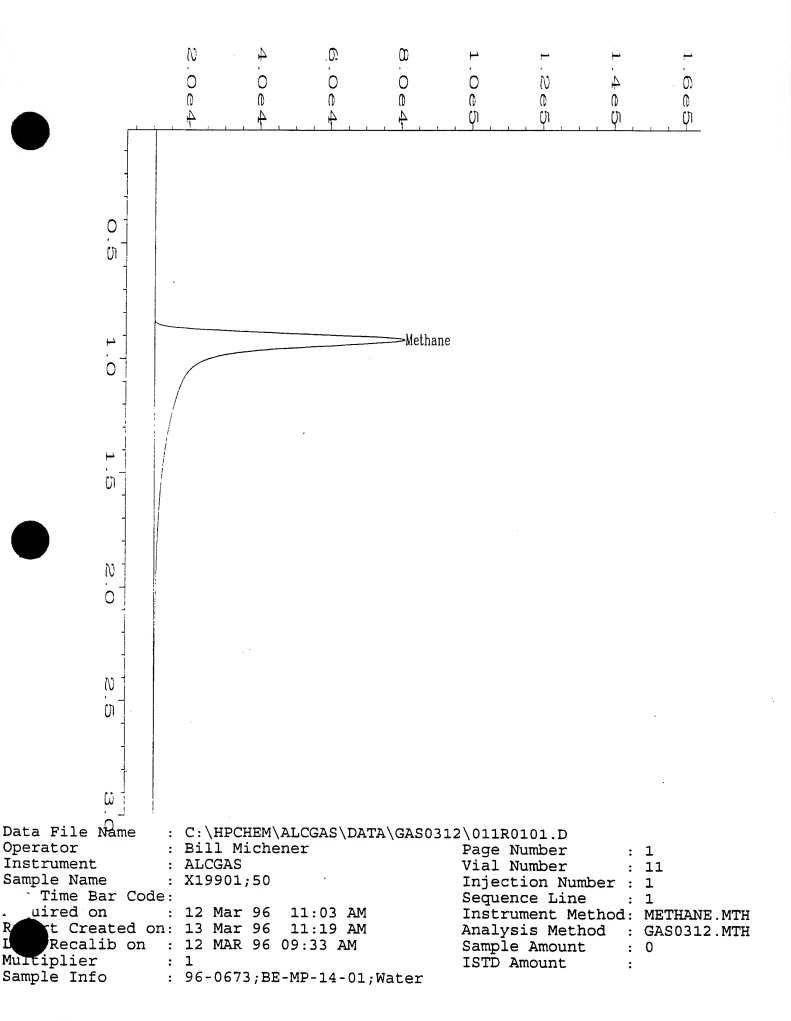
B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Analyst

Approved



EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP-26-01	Client Project No.	: 722450.14020 Bea
Lab Sample Number	: X19902	Lab Project No.	: 96-0673
Date Sampled	: 3/6/96	Dilution Factor	: 1.00
Date Received	: 3/8/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/12/96	Matrix	: Water
Date Analyzed	: 3/12/96	Lab File No.	: GAS0312012

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L
Methane	74-82-8	U	0.002

Temperature Amount Injected Total Volume of Sample Head space created Methane Area		72.6 F 0.5 ml 43 ml 4 ml 0 ug	Saturation Concentration Concentration in Head Space	Meth Meth	<u> </u>
--	--	-------------------------------	---	--------------	----------

16 g

QUALIFIERS:

E = Extrapolated value.

Atomic weight(Methane)

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Analyst

K Hollman
Approved

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Operator
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Sample Name
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                                                 Instrument Method: METHANE.MTH
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                                                 Analysis Method
                                                                   : GAS0312.MTH
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                                                 Sample Amount
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                                                 ISTD Amount
Sample Info
                 : 96-0673;BE-MP-27-01;Water
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EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP-14-01	Client Project No.	: 722450.14020
Lab Sample Number	: X19901	Lab Project No.	: 96-0673
Date Sampled	: 3/6/96	Dilution Factor	: 50.00
Date Received	: 3/8/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/12/96	Matrix	: Water
Date Analyzed	: 3/12/96	Lab File No.	: GAS0312011

Compound Name	Sample Cas Number Concentration mg/L		RL mg/L	
Methane	74-82-8	3.2	0.1	

Temperature	:	72.3 F	Saturation	Meth	0.77883 ⁻ 1
Amount Injected	•	0.01 ml	Concentration	•	
Total Volume of Sample	:	43 ml	Concentration	Meth	2.452556
Head space created	:	4 ml	in Head Space	•	
Methane Area	:	362.229 ug			

Atomic weight(Methane) : _____ 16 g

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Analyst

Approved

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Instrument Vial Number 2 Sample Name : GB031296 Injection Number : Time Bar Code: Sequence Line : 1 : 12 Mar 96 sired on 08:57 AM Instrument Method: METHANE.MTH t Created on: 12 Mar 96 09:41 AM Analysis Method : GAS0312.MTH Recalib on : 12 MAR 96 09:33 AM Sample Amount : 0 Multiplier : 1 ISTD Amount : Sample Info : Gas Method Blank

Operator

Displaced 4ml of deionized water in 43ml vial with Helium, shook for 5 min and injected 500ml

EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form

Client Sample Number	: BE-MP-27-01	Client Project No.	: 722450.14020 Bea
Lab Sample Number	: X19900	Lab Project No.	: 96-0673
Date Sampled	: 3/6/96	Dilution Factor	: 1.00
Date Received	: 3/8/96	Method	: RSKSOP-175
Date Extracted/Prepared	: 3/12/96	Matrix	: Water
Date Analyzed	: 3/12/96	Lab File No.	: GAS0312010

Compound Name	Cas Number	Sample Concentration mg/L	RL mg/L	
Methane	74-82-8	U	0.002	

Temperature	:	70.6 F	Saturation	Meth	n
Amount Injected	:	0.5 ml	Concentration		-
Total Volume of Sample	:	43 ml	Concentration	Meth	
Head space created	:	4 ml	in Head Space		
Methane Area	:	0 ug			

Atomic weight(Methane) : _____ g

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

Analyst

K. Hollman
Approved

AF0673.XLS

EVERGREEN ANALYTICAL, Inc.

4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Total Organic Carbon

Date Sampled : 3/06-07/96 Date Received : 3/08/96

Client Project ID.

: 722450.14020 Beale

Date Prepared

Lab Project Number: 96-0673

Date Analyzed

: 3/13/96 : 3/13/96 Method

: EPA 415.1

Detection Limit

: 1.0 mg/L

Evergreen Sample #	Client Sample ID.	<u>Matrix</u>	Total Organ <u>Carbon</u>	nic	Dilution <u>Factor</u>
X19900	BE-MP-27-01	Water	<1.0	mg/L	1
X19901	BE-MP-14-01	Water	17.6	mg/L	1
X19902	BE-MP-26-01	Water	<1.0	mg/L	1
X19903	BE-MP-24-01	Water	<1.0	mg/L	1
X19904	BE-MP-03-01	Water	1.4	mg/L	1
X19905	BE-MP-25-01	Water	<1.0	mg/L	1
Method Blank	(3/13/96)	Water	<1.0	mg/L	

Quality Assurance

		Spike Amount (mg/L)	Sample Result (mg/L)	Spike Result (mg/L)	% Recovery
X19900	Matrix Spike BE-MP-27-01	10.0	<1.0	10.2	102
X19900	Matrix Spike Du BE-MP-27-01	ıp 10.0	<1.0	10.1	101
MS/MSD	RPD				0.79

Approved

EVERGREEN ANALYTICAL, INC. 4036 Youngfield St. Wheat Ridge, CO 80033 (303) 425-6021

Methane Report Form Method Blank Report

Method Blank Number

: GB031296

Client Project No.

: 722450.14020 Beal

Date Extracted/Prepared

: 3/12/96

Lab Project No.

: 96-0673

Date Analyzed

: 3/12/96

Dilution Factor

: 1.00

Method

Matrix

: RSKSOP-175

: Water

Lab File No.

: GAS0312002

		Sample		
Compound Name	Cas Number	Concentration	RL .	
		mg/L	mg/L	
Methane	74-82-8	U	0.002	

QUALIFIERS:

E = Extrapolated value.

U = Compound analyzed for, but not detected.

B = Compound also found in the blank.

RL = Reporting Limit.

NA = Not Available/Not Applicable.

L. Hollman **Approved**

AF0673.XLS

TURNAROUND REQUIRED* 🗡 STD (2 wks) 🗀 STD UST (3 day) 96-0673 PROJECT ID 722450, 14020 Bec/e in shaded area PO.# 722450 EAL Sample No. EAL use only Do not write 9400 903 900 402 9 Custodian Other (Specify). *expedited turnaround subject to additional fee CLIENT CONTACT (print) W (CLIC, e **JALYTICAL SERVICES REQUEST** ग्राध EAL. QUOTE # **ANALYSIS REQUESTED** 705 TEPH 8015mod. (Diesel) Wheat Ridge, Colorado 80033 (303) 425-6021 FAX RESULTS Y / (N_) Evergreen Analytical Inc. DOD XXI FAX (303) 425-6854 4036 Youngfield St. BTEX 8020/602 (circle)/MTBE (circle) (800) 845-7400 Herbicides 8150/515 (circle) FAX# 510 7699244 CHAIN OF CUSTODY RECO WOULD JULY OKNY STED アイションコン **MATRIX** MICHAEL ZIP CASE C. 4 1A25 8 TIME 200 2 PARSONSES SAMPLED 1796 0000 BE MP 26-01 B/6 196 BE- N. P. 24-01 3/6/96 3/6/96 BE-WRO3-01/3/1/96 DATE CITY ALANCO ASTATE CA Please PRIN all information: 3 Evergreen Analytical Cooler No. Sampler Name: 53 BE-IMP25-01 BE-MP-27-01 BE-MP-14-01 **IDENTIFICATION** (print) M. chrel PHONE SIC SAMPLE CLIENT Cooler Received COMPANY ADDRESS

3/16/Time Received by: (Signature)

Date/Time Relinquished by: (Signature)

JAME WALKS CONSTINE Date/Time Re

Container Size

TO SOLVE IN

37.8

Location

Relinquished by: (Signature)

Instructions:

9 Ϊ

•	nalytical Sa		eceipt			
Date & Time Rec'd: 3		35 s	hipped	v1a:	1-X	
Client: Parson			···	(Airbill #	11540	92763
Client Project ID(s):	722450	1.1407	0 B	eole.		
EAL Project #(s):96-	0673		EAL	Cooler(s)	· (Y)	N
Cooler# 730	· · · · · · · · · · · · · · · · · · ·					
Ice packs (Y) N	Y N	Y	N	y N	Y N	
Temperature ℃				_		-
				Y	N	N/A
1. Custody seal(s) pro Seals on cooler : Seals on bottle :	intact			X		X
2. Chain of Custody pr	resent:			X_		
3. Samples Radioactive	Comment on COC if	> 0.5mr/h)			_X	
4. Containers broken o	or leaking: c	omment on CO	C if Y)			
5. Containers labeled:				<u>X</u>		
6. COC agrees w/ bottl	es received:	(Comment o	a COC if N	<u>X</u>		
7. COC agrees w/ label	S: (Comment on COC	if N)		X		
8. Headspace in vials-	waters only:	(Comment	n COC if Y		<u> </u>	
9. VOA samples preserv	ed: Mit	thones.	•	X		
10. pH measured on metal List discrepancies *Non-EAL provided conta						<u> </u>
11. Metal samples prese Total, Disso D or PD to be filte	ent: olved				<u> </u>	
T,TR,D,PD to be Pre						
12. Short holding times Specify parameters		UZ				
13. Multi-phase sample(s) present:				X	
14. COC signed w/ date/	time:					
Comments:						
(Additional comments on back		\				
Custodian Signature/Dat		Mm	3/8	196		
J / Dau				- L C -		

Evergreen Analytical Sample Log Sheet	Project # <u>96-0673</u>
Dat (s) Sampled: 03/06,07/96 COC	Date Due: 03/22/96
te Received: 03/08/96 0935 nt Project I.D. 722450.14020 BEAT	Holding Time(s): 03/08,09-NO ₂ ,NO ₃ 3/20,21-ALKALINITY Rush STANDARD
Client: PARSONS ENGINEERING SCIENCE, Address: 1301 MARINA VILLAGE PKWY #200 ALAMEDA, CA 94501 Contact: MICHAEL PHELPS Phone #510-769-0100 Fax #510-769-9244 Special Invoicing/Billing	E.A. Cooler # 720 Airbill # FEDEX 1154092763 Client P.O. 722450
Special Instructions	
Lab Client ID # ID# Analy	sis Mtx Btl Loc
X19900A BE-MP-27-01 TOO	W 125A C2
X19901A BE-MP-14-01 TOO	
9902A BE-MP-26-01 TOO	
9903A BE-MP-24-01 TOO	
04A BE-MP-03-01 TOO	
X19905A BE-MP-25-01 TOO	
X19900B-D BE-MP-27-01 METHA	
X19901B-D BE-MP-14-01 METHA	
X19902B-D BE-MP-26-01 METHA	
X19903B-D BE-MP-24-01 METHA	
X19904B-D BE-MP-03-01 METHA	
X19905B-D BE-MP-25-01 METHA	NE W 40V 2
X19900E BE-MP-27-01 AMMON	IA W 500P C2
X19901E BE-MP-14-01 AMMON	IA W 500P C2
X19902E BE-MP-26-01 AMMON	IA W 500P C2
X19903E BE-MP-24-01 AMMON	IA W 500P C2
X19904E BE-MP-03-01 AMMON	IA W 500P C2
X19905E BE-MP-25-01 AMMON	IA W 500P C2
R=Sample to be returned	
ite GC/MS GC X Metals	Wet Chem X HPLC SxPrep
SxRec C QA/QC C	Acctg C File Orig
Page 1 of 2 Page(s)	Custodian/Date: MM/PLB5/8/16

Lab ID #	Client ID#	Analysis	Mtx	Btl	Loc
X19900F	BE-MP-27-01	ALKALINITY	W	125P	C2
X19901F	BE-MP-14-01	ALĶALINITY	W	125P	C2
X19902F	BE-MP-26-01	ALKALINITY	W	125P	C2
X19903F	BE-MP-24-01	ALKALINITY	W	125P	C2
X19904F	BE-MP-03-01	ALKALINITY	W	125P	C2
X19905F	BE-MP-25-01	ALKALINITY	W	125P	C2
X19900G	BE-MP-27-01	NO ₂ , NO ₃ , SO ₄ , Cl ⁻	W	40V	2
X19901G	BE-MP-14-01	NO ₂ , NO ₃ , SO ₄ , C1	W	40V	2
X19902G	BE-MP-26-01	NO ₂ , NO ₃ , SO ₄ , Cl ⁻	W	40V	2
X19903G	BE-MP-24-01	NO ₂ , NO ₃ , SO ₄ , Cl ⁻	W	4 0 V	2
X19904G	BE-MP-03-01	NO ₂ , NO ₃ , SO ₄ , Cl ⁻	W	40V	2
X19905G	BE-MP-25-01	NO ₂ , NO ₃ , SO ₄ , Cl ⁻	W	40V	2

Page 2 of 2 Pages
Project # 96-0673

R=Sample to be returned



Hazen Research, Inc.

4601 Indiana St. • Golden, Colo. 80403 Tel: (303) 279-4501 • Telex 45-860

FAX: (303) 278-1528

Date HRI Project 009-93 HRI Series No. C038/96 Date Rec'd. 03/01/96 Cust. P.O. #

March 7 1996 03/01/96

Evergreen Analytical Inc. Carl Smits 4036 Youngfield Wheat Ridge CO 80033

SCREEN ANALYSIS REPORT

Sample Number: C038/96-1

Sample Identification: 96-0587 BE-MP24-3 02/26/96 @ 1110

	· · ·				
Mesh Size Pass-Retair		Direct Weight,g	Direct Weight,%	Cum Weight % Retained	Cum Weight % Pass
- 4 4 - 6 6 - 8 8 - 10 10 - 14 14 - 20 20 - 28 28 - 35 35 - 48 48 - 65 65 - 100 100 - 150 150 - 200 200 - 270 270 - 325 325 - PAN	Tota1	14.1 4.4 5.2 3.8 4.6 4.6 5.7 7.5 8.7 11.0 11.4 15.3 3.6 141.2 254.2	5.55 1.73 2.04 1.26 1.89 1.81 2.24 2.95 3.50 3.42 4.33 4.48 6.02 1.42 55.55 100.00	5.55 7.28 9.32 10.58 12.47 14.28 16.09 18.33 21.28 24.78 28.20 32.53 37.01 43.03 44.45 100.00	94.45 92.72 90.68 89.42 87.53 85.72 83.91 81.67 78.72 75.22 71.80 67.47 62.99 56.97 55.55 0.00

By:

Robert Rostad

Laboratory Manager

Screen sizes are in Tyler Mesh.

Project # %-0869, 0587

& MOISTURE

	Original														-									-		
$\frac{(A-B)}{C} \times 100$	& Wofature		18.06	15.63	7.49	52.61	21.59	7.74	1421	6.58	6.62															
υ	Net Wt. of Wet Sample	16.04	14.95	10/1	/3.34	1372	11.39	19.61	14.00	15.18	15.09															
Ф	Tin + Sample Drv	14.36	1341	h6.61	12.61	12.15	90.01	8/1:1/	13.15	jų.id	14 43															
A	Tin + Sample	17.16	// · 9/	15.13	14.48	14.86	12.52	14.04	15.14	1627	16.33															
3/32/46	pH Tin	1.12	1.16	1/./3	1/1/	h/·/	1/3	1.13	1 /1// 1	1 60 / 1	1.13															
		COG7 1×20886	2 887	3 888	4 889	5 890	168 9	7 893		03x1x19543	10 543	11	12	13	14	15	16	17	18	19	20	21	22	23	24	

HUFFMAN

LABORATORIES, INC.

Quality Analytical Services Since 1936 4630 Indiana Street • Golden, CO 80403

ANALYSIS	CARBONATE	METHOD	SOP COU-02
ANALYZER#	6	COULOMETER#	Z
BALANCE#	10		

j	ALCIUM CA	ARBONATE CaCO₃	BOTTLE # 177/2	% C THEO	RY = 12.00%	SODIUM CARBO (STD # 134) Na ₂		BOTTLE# 730+Y7/3	%C THEOR	RY = 11.33 %
	MPLE NO.	TARE WT. GRAMS	TARE + SAMPLE WT.	SAMPLE WT. GRAMS	NOTES	(COUNTS) μ GRAMS CARBON	LESS BLANK	% CARBONATE CARBON AS CARBON	QC	% RECOVERY
\int	125601	0.896936	1.78334/	0.836405	VIII)	(1357.9)	13509			
	133	0;877370	0.888844	0.011474	sp.kg/					
S	125601	0.888038	1.757185	0.869147		(1310.9)	1303.9			
(133	0.4/1120	0.922427	0.011307	Spike/	1	1240.6	11.90-X	3.7	00
	/33	0.928347	0.938 767	0.010420		1247.6	£47. J-	11.90. ×-	CCS	99.2
			7060 7							t0.7%.d
	122	A 0051 1/2	C 0 - 1 - C/2	PELKUV4		1244.9	1237.9	11.85	(CS	98.8
-	/.33	0.895693	0.906087	4.010911		17-11. /	.,,	111.00		75. 8
-										
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	YST //	?lle	DATE	7-96	REVIEW	, <u>, ,</u>	DAT	3/8/96	PAGE 2	OF Z



Hazen Research, Inc.

4601 Indiana St. • Golden, Colo. 80403 Tel: (303) 279-4501 • Telex 45-860 FAX: (303) 278-1528 HRI Project 009-93 HRI Series No. C038/96 Date Rec'd. 03/01/96 Cust. P.O. #

Date

March 7 1996 009-93 C038/96 03/01/96

Evergreen Analytical Inc. Carl Smits 4036 Youngfield Wheat Ridge CO 80033

SCREEN ANALYSIS REPORT

Sample Number: C038/96-2

Sample Identification: 96-0587 BE-MP25-7 02/28/96 @ 1040

Mesh Size Pass-Retained	Direct Weight,g	Direct Weight,%	Cum Weight % Retained	Cum Weight % Pass
- 4 4 - 6 6 - 8 8 - 10 10 - 14 14 - 20 20 - 28 28 - 35 35 - 48 48 - 65 65 - 100 100 - 150 150 - 200 270 - 270 270 - 325 325 - PAN	64.8 16.3 14.8 14.6 18.0 19.3 21.2 20.7 15.1 7.5 4.0 3.5 2.8 2.7 0.5 38.0	24.56 6.18 5.61 5.54 6.82 7.32 8.04 7.85 5.72 2.84 1.52 1.33 1.06 1.02 0.19 14.40 100.00	24.56 30.74 36.35 41.89 48.71 56.03 64.07 71.92 77.64 80.48 82.00 83.33 84.39 85.41 85.60 100.00	75.44 69.26 63.65 58.11 51.29 43.97 35.93 28.08 22.36 19.52 18.00 16.67 15.61 14.59 14.40 0.00

By:

Robert Rostad

Laboratory Manager

Screen sizes are in Tyler Mesh.



LABORATORIES, INC.

Quality Analytical Services Since 1936
4630 Indiana Street • Colden, CO 80403

ANALYSIS	CARBONATE CARBON	METHOD	SOP COU-02
ANALYZER#	6	COULOMETER#	ユ
BALANCE#	1/2		

ALCIUM C	ARBONATE) CaCO ₃	BOTTLE# 43/2	% C THEO	RY = 12.00%	SODIUM CARBO (STD # 134) Na ₂		30TTLE# <i>33014713</i>	%C THEO	RY = 11.33 %
MPLE NO.	TARE WT. GRAMS	TARE + SAMPLE WT.	SAMPLE WT. GRAMS	NOTES	(COUNTS) μ GRAMS CARBON	LESS BLANK	% CARBONATE CARBON AS CARBON	QC	% RECOVER
B/					8.0			IB	
B/					7.0			MB	
					1 - 7				
133	C.877466	0.888050	0010584		1272.6	1265.6	11.96	ICS	99. C
133		C896070			1299.9	12929	11.92	(15	99.3
134	1	0.87776 C			1157.2	1150.2	1124	LCS	99.2
•			7, 20						
						226.5		V • 1	
25601	6.855974	1502674	0.646700		8.8	1.8	21.01		
601	0.898797	1.925798	1.026999 0. ×m 3.5	<u></u>	7.4 .	0.4	60.01		
25602	0.862578	1.776263	0.913685		7.4	0.4	.20.01		
	0.005782	1.6_1717	0.757763				7		
133	0.882893	0.893472	6.010579		12776	1270.6	12-01	(<i>(</i> S	100.1
									
		121e	needs	dup /5	rikes		-	<u> </u>	
				·					
107		- I DAT	F	REVIEWE	D	DATE			,
VSI /	Elli-	3.	E 5 -96	KCAIEAAE	RUK		030596	PAGE /	OF / ED 7/13/95

HUFFMAN

LABORATORIES, INC.

Quality Analytical Services Since 1936 4630 Indiana Street - Golden, CO 80403

ANALYSIS	CARBONATE CARBON	METHOD	SOP COU-02
ANALYZER#	6	COULOMETER#	2
BALANCE #	10		

CALCIUM CA (STD # 133)	ARBONATE CaCO ₃	BOTTLE# 4712	% C THEOR	RY = 12.00%	SODIUM CARBO (STD # 134) Na ₂		30TTLE# 273011913	%C THEO	RY = 11.33 %
SAMPLE NO.	TARE WT.	TARE + SAMPLE WT.	SAMPLE WT. GRAMS	NOTES	(COUNTS) µ GRAMS CARBON	LESS BLANK	% CARBONATE CARBON AS CARBON	QC	% RECOVERY
81					7.2			IB	
B1					7.6			MB	
					7.0				
133	C 4/, U/ 32	0874654	0.009981		1198 4	1191.4	11.94	Ics	99.5
122		0.892334				1209.4	11.88	((5	99.0
. 211		0.891505				1168.6	11.32	LCS	99.9
	0.881187	0.897.305							
		-00)							
						736 =	A-4-5	<u> </u>	
13477				//	10 8m3-7	1/2	//-		
		0970192							
			. (0)/02				% C	201	Mesa
			07,7007		7	·//:	· · · · · · · · · · · · · · · · · · ·		+000% of
			.416013		105.1	., 0.,			
		70))	181 020.		10	,	7. 10		
33	901715	0.912368	0.010653		1279.6	272.6	11.95	(15	
1/	2 872-07								
ALYST ATA			E 7-96	REVIEWE			18/96		0F 2



Quality Analytical Services Since 1936
4630 Indiana Street • Colden, CO 80403

ANALYSIS: TOTAL CARBON	METHOD: HIGH TEMP COMB INFRARED DET.
INSTRUMENT : LECO CR12	ANALYZER # 7
BALANCE# 19	

D. CAL JFFMAI 12.00 %C	CIUM CARBONAT N std # 133) (theory)	е ви_23	140	(HUFFN	I.S.T. BUMAN std # C (theory			MENT 27 485	
SAMPLE #	SAMPLE WT G			CAF	% RBON RE- ALIB	% CARBON POST- CALIB		QC	% REC.
10)					·				
				70	.1 /				
					10				
/33	0.0980			12.		11.99		Tcs	99.9
133	0.0970					11.96		ccs	99.8
. ;									
127	D 09G D					12.06		(cs	100.5
133 B1	1.0000		44-45-0			0,003		TB	700.5
BI	1.0000					0.008		MB	
W2704	0.2549		- Marine			3 3.32		LCS	99.5
									7
LYST	DATE 3-12	7 - 9.6	REVIEWED	55	DATE	3/12/96	PAC	GE .	OF 3



LABORATORIES, INC.

Quality Analytical Services Since 1936 4630 Indiana Street • Golden, CO 80403

ANALYSIS: TOTAL CARBON METHOD: HIGH TEMP COMB. - INFRARED DET. INSTRUMENT: LECO CR12 ANALYZER # 7

			BALANCE	# 19)			
STD. CALC (HUFFMAN 12.00 %C (1		ATE BN <u>2</u>	7340	HL (HL	D. N.I.S.T. B JFFMAN std : 48 %C (theor		MENT 27	
SAMPLE #	SAMPLE WT G				% CARBON PRE- CALIB	% CARBON POST- CALIB	QС	% REC.
133	0.1/70					12.03	ccs	100.3
	=	-	0,0					#1.Co
125601	0.9900	7	(Entered)			(1.416)	MS	
133	0.1030	5	Spiko				MS	
125601	0.9740	2	(Entered)			(1.413)	MSD	
133	0.1010)	Spike				MSD	
N2704	0.3090					3.334	(CS	99.6
							 :	
				_				

ANALYSI .	DATE 3-12-96	REVIEWED 5.	DATE 3/12/96	PAGE OF	
	()		P /	rev 7/3/95	5



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LABORATORIES, INC.

Quality Analytical Services Since 1936
4630 Indiana Street • Colden, CO 80403

ANALYSIS: TOTAL CARBON	METHOD: HIGH TEMP
7.17.12.10.10.1	COMB INFRARED DET.
INSTRUMENT: LECO CR12	ANALYZER # 7
BALANCE# 19	

STD. N.I.S.T. BUFFALO RIVER SEDIMENT 2704 D. CALCIUM CARBONATE (HUFFMAN std # N2704) FFMAN std # 133) BN 485/ 3.348 %C (theory) BN 2340 172.00 %C (theory) % CARBON % CARBON SAMPLE SAMPLE WT REC. QC POST-PRE-CALIB CALIB 100.3 Ics 12.04 133 0-1070 11.84 78. X 65 0.1180 133 TB BI 0.002 1,0000 MB 131 0.005 1.0000 99.4 3.329 LCS 102704 0.3040 (15 12.19 101.6 0.1080 REVIEWED C DATE / 3/12/96 PAGE DATE 3-4-96 OF 2

rev 7/3/95



LABORATORIES, INC.

Quality Analytical Services Since 1936 4630 Indiana Street • Golden, CO 80403

ANALYSIS: TOTAL CARBON	METHOD: HIGH TEMP COMB INFRARED DET.
INSTRUMENT: LECO CR12	ANALYZER# 7
DALANCE # 10	

STD. N.I.S.T. BUFFALO RIVER SEDIMENT 2704 STD. CALCIUM CARBONATE (HUFFMAN std # N2704) (HUFFMAN std # 133) BN 4851 BN 234 0 3.348 %C (theory) 12.00 %C (theory) % CARBON % CARBON SAMPLE WT SAMPLE QC REC. POST-PRE-G CALIB CALIB 0.205 0.6010 125601 WH) £ 2.0%. DUP 0.197 125601 0.4870 of May 0.032 0.9640 125602 (15 101.2 12.20 133 0.1190

REVIEWED S PAGE DATE 3.4-96 rev 7/3/95



Quality Analytical Services Since 1936 4630 Indiana Street • Golden, CO 80403

NON-CLP ANALYSIS RESULTS ABORATORY CONTROL STANDARD

ےate:

03/13/96

Client: Evergreen Analytical

Lab Name:

Huffman Labs

Contact: Patty McClellan

Contact:

Sue Zeller

Huffman Lab #: 125696

LABORATORY CONTROL STANDARD

Lab	Source	Element/	True	Found	% R	Units		Method	Instrument
· ID#		Compound	Value	Value			Date	#	ID
LCS	BN 4851	TC	3.35	3.33	99	%	03/04/96	Le∞ CR12	#7
LCS	BN 4851	TC	3.35	3.33	99	%	03/12/96	Leco CR12	#7
LCS	BN 2703	CC	11.33	11.24	99	%	03/05/96	COU-02	#2
LCS	BN 2703	CC	11.33	11.32	100	%	03/07/96	COU-02	#2

SPIKE RECOVERY

Lab	Source	Element/	True	Found	% R	Units		Method	Instrument
ID#		Compound	Value	Value			Date	#	ID
SPIKE	BN 4712	TC	12360	11988	97	ug C	03/12/96	Leco CR12	#7
SPIKE DUP	BN 4712	TC	12120	13662	113	ug C	03/12/96	Leco CR12	#7
SPIKE	BN 4712	CC	1377	1351	98	ug C	03/07/96	COU-02	#2
SPIKE DUP	BN 4712	CC	1357	1304	96	ug C	03/07/96	COU-02	#2



Quality Analytical Services Since 1936 4630 Indiana Street • Golden, CO 80403

NON-CLP QA/QC ANALYSIS RESULTS INITIAL AND CONTINUING CALIBRATION VERIFICATION

Date:

03/13/96

Client: Evergreen Analytical

Lab Name:

Huffman Labs

Contact: Patty McClellan

Contact:

Sue Zeller

Huffman Lab #: 125696

INITIAL CALIBRATION

	Lab	Source	Element/	True	Found	% R	Units		Method	Instrument
	ID#		Compound	Value	Value			Date	#	ID
	ICS	BN 4712	TC	12.00	12.04	100	%	03/04/96	Leco CR12	#7
•	ICS	BN 4712	TC	12.00	11.99	100	%	03/12/96	Leco CR12	#7
	ICS	BN 4712	CC	12.00	11.96	100	%	03/05/96	COU-02	#2
	ICS	BN 4712	CC	12.00	11.94	100	%	03/07/96	COU-02	#2

Slope =

NA

Intercept =

NA

Single point calibrations for this test.

95% Correlation Coefficient =

NA

CONTINUING CALIBRATION VERIFICATION

Instrument	Method		Units	% R	Found	True	Element/	Source	Lab
# ID	#	Date			Value	Value	Compound		ID #
2 #7	Leco CR12	03/04/96	%	102	12.19	12.00	TC	BN 4712	CCS
2 #7	Leco CR12	03/04/96	%	102	12.20	12.00	TC	BN 4712	CCS
	Leco CR12	03/12/96	%	100	12.03	12.00	TC	BN 4712	CCS
	Leco CR12	03/12/96	%	99	3.33	3.35	TC	BN 4851	CCS
	COU-02	03/05/96	%	99	11.92	12.00	CC	BN 4712	CCS
· · ·	COU-02	03/05/96	%	100	12.01	12.00	CC	BN 4712	CCS
	COU-02	03/07/96	%	100	11.95	12.00	CC	BN 4712	CCS
1.2	COU-02	03/07/96	%	99	11.91	12.00	CC	BN 4712	ccs

Evergreen Analytical Sample Receipt	Check-in Record
Date & Time Rec'd: 3/1/96 /000 Shipped	Via: Feder
client: Parsons ES	(Airbill # if applicable)
Client Project ID(s): 722450, 14020	<u> </u>
0. — 10	Ccoler(s): Y N
cooler# Che, L	
Ice packs Y N Y N Y N	y n y n
Temperature °C O	
	Y N N/A
<pre>1. Custody seal(s) present: Seals on cooler intact</pre>	
Seals on bottle intact	
2. Chain of Custody present:	*
3. Samples Radioactive: (Comment on COC if > 0.5mr/h)	<u>~</u> ×
	<u> </u>
4. Containers broken or leaking: (Comment on COC if Y)	
5. Containers labeled:	
6. COC agrees w/ bottles received: (Comment on COC if N)	
7. COC agrees w/ labels: (Comment on COC if N)	<u> </u>
8. Headspace in vials-waters only: (Comment on COC If Y)	<u> </u>
9. VOA samples preserved:	
10. pH measured on metals, cyanide or phenolic	s*:
List discrepancies*Non-EAL provided containers only, water sample	og on?w
·	es only.
11. Metal samples present:	
Total, Dissolved, TCLP D or PD to be filtered:	
T,TR,D,PD to be Preserved:	
12. Short holding times:	×
Specify parameters	
13. Multi-phase sample(s) present:	
14. COC signed w/ date/time:	
Comments:	· · · · · · · · · · · · · · · · · · ·
Additional comments on back)	· · · · · · · · · · · · · · · · · · ·
Custodian Signature/Date:	96



Quality Analytical Services Since 1936 4630 Indiana Street • Golden, CO 80403

NON-CLP ANALYSIS RESULTS

Date:

03/13/96

Lab Name:

Huffman Labs

Sue Zeller

Client: Evergreen Analytical

Contact: Sample Matrix:

soil

Contact: Patty McClellan

Huffman Lab #: 125696

	Client Smp#	Lab ID#	Element/ Compound	Dilution Factor	Results	Units	Prep Date	Analysis Date	Sample Size (g)	Method #	Instrument
	BE-MP24-3	12569601	TC	NA	0.21	%	NA	03/04/96	0.601	Leco CR12	#7
	BE-MP24-3	12569601	TC	NA	0.20	%	NA	03/04/96	0.987	Leco CR12	#7
	BE-MP25-7	12569602	TC	NA	< 0.05	%	NA	03/04/96	0.964	Leco CR12	#7
	BE-MP24-3		CC	NA	< 0.02	%	NA	03/05/96	0.647	COU-02	#2
	BE-MP24-3		CC	NA	< 0.02	%	NA	03/05/96	1.027	COU-02	#2
	BE-MP25-7	12569602	CC	NA	< 0.02	%	NA	03/05/96	0.913	COU-02	#2
		% mc	isture		adjust	ed result					
X19542	BE-MP24-3	12569601	6.58 TOC	NA	0.21), az %	NA	NA	NA	by calc	NA
X19542	BE-MP24-3	12569601	6.58 TOC	NA	0.206	2.21 %	NA	NA	NA	by calc	NA
X19543	BE-MP25-7	12569602	6.62 TOC	NA	< 0.05	0.05%	NA	NA	NA	by calc	NA

Samples analyzed and results reported on as as received basis.

Soil samples are not homogeneous.

Values reported below Detection Limits are for reference only.

TC detection limit = 0.05%

CC detection limit = 0.02%

TOC detection limit = 0.05%

Evergreen Analytical Sample Log Sheet	Project # <u>9</u> 5-0587
Date(s) Sampled: 02/26,28/96 COC	Date Due: 03/15/96
e Received: 03/01/96 1000	Holding Time(s):
roject I.D. 722450.14020	Rush STANDARD
Client: PARSONS ENGINEERING SCIENCE, INC. Address: 1301 MARINA VILLAGE PKWY #200 ALAMEDA, CA 94501 Contact: MICHAEL PHELPS Phone #510-769-0100 Fax #510-769-0100 Special Invoicing/Billing	Cooler Return 5.00 E.A. Cooler # N/A Airbill # FEDEX 1154092542 Client P.O. 722450
Special Instructions	
Lab Client ID # ID# Analysis	Mtx Btl Loc
X19542A BE-MP24-3 SIEVE ANALYSI X19543A BE-MP25-7 SIEVE ANALYSI	
R=Sample to be returned Le GC/MS GC Metals W SxRec C QA/QC C Accord Page 1 of 1 Page(s)	•

CHAIN OF CUSTODY RECORD / ANALYTICAL SERVICES REQUEST





nature)

Case Narrative Parsons Engineering Science, Inc. Page Two

General Chemistry

There are no quality control anomalies to report for the following analyses: Ammonia by Standard Method 4500-NH₃F, Alkalinity by EPA Method 310.1, Total Organic Carbon in water by EPA Method 415.1 or Anions by EPA Method 300.0.

Total Organic Carbon in soil

Total Organic Carbon (TOC) in soil was analyzed by Huffman Laboratories of Golden, Colorado. TOC is determined by analyzing for total carbon (TC) and inorganic (carbonate) carbon (CC), then calculating the difference as TOC. The reports from Huffman are included.

Grain Size

Grain size is determined by sieve analysis as performed by Hazen Research Laboratory of Golden Colorado. The report from Hazen is included. Due to the nature of this report, it is not included on the electronic diskette.

Disk Deliverables

The disk deliverables are included with the hard copy data package. MS/MSD and laboratory duplicate samples are not included on the disk. Please note that blank spaces in the laboratory detection limit and/or practical quantitation limit (PQL) column indicate that there is no detection limit or PQL for that analyte.

Reporting limits have been adjusted to reflect percent moisture content in all soil samples or increases due to dilutions.

A hard copy of each spreadsheet included on the diskette is included. The name for each spreadsheet is located in the top left corner on the first page of each printout.

Patricia A. McClellan, Program Manager



April 1, 1996

MR MICHAEL PHELPS PARSONS ENGINEERING SCIENCE INC 1301 MARINA VILLAGE PKWY #200 ALAMEDA, CA 94501

Data Report: 96-0587,0673,0686,0706,

0741,0776,0805

Client Project: 722450.14020 BEALE AFB

Dear Mr. Phelps:

Enclosed are the analytical results for the samples shown in the Sample Log Sheet. The enclosed data have been reviewed for quality assurance. If you have any questions concerning the reported information, please contact Patty McClellan, Program Manager.

Upon completion of all required analyses and acceptance of the data report by PES (within 3 weeks of final data package deliver), EAL will be responsible for proper disposal of any remaining samples, sample containers, shipping containers, and Sytrofoam or plastic packing materails in accordance with sound environmental practices, based on the sample analytical results. However, EAL will give prior notification to and receive the approval of PES before disposing of any remaining samples. EAL will maintain proper records of waste disposal methods and disposal methods and disposal company contracts on file for inspection.

The invoice for this work will be mailed to your Accounts Payable department shortly.

Thank you for using the services of Evergreen Analytical.

Sincerely,

Jack Barney

President

JB/mp



CASE NARRATIVE

Evergreen Analytical Laboratory (EAL) Projects: 96-0587, 96-0673 96-0686, 96-0706, 96-0741, 96-0776, 96-0805

Parsons Engineering Science, Inc. (PES) Project:

Beale AFB 722450.14020

Sample Receipt

Between March 1 and March 16, 1996, soil and groundwater samples were received at EAL for analysis under subcontract 722450.SC02. Refer to the EAL Check-in Record for specific information regarding the condition of samples upon receipt. Refer to the EAL Sample Log Sheet for specific log-in information and cross-reference of EAL and PES sample identifications.

Data Package

All data are reported in one comprehensive package that is segregated based upon EAL project number. Each EAL project represents a group of samples received on a given day. The EAL Sample Log Sheet summarizes the samples represented in each EAL project.

A separate invoice for each EAL Project number will be generated.

Quality assurance data may overlap from one EAL project to another. All required matrix spike/matrix spike duplicate (MS/MSD) and laboratory duplicate samples were analyzed when required and also are included in the data package.

Methane, Method RSKSOP-175

All samples submitted for methane analysis were analyzed within holding time.

The spike recovery in the Laboratory Control Sample LCS031296 was above the acceptance limits of 67 - 85 percent, at 93 percent. This spike recovery is considered to be acceptable.

There are no other quality control anomalies to report.

atch' N	0.: /	/ <i>HCL</i> Date: <u>5/21/96</u> 27723		fix: 142 JKL	Co	Page ntinued from Page:	65 64
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Curtis & Tompkins, Ltd. Sample Batch Report

Batch Number: 27719 Date Started: 21-MAY-96 Batched By : Anna Maria Pajarillo

Analysis : N/A Bgroup: : TVH Department: Volatile Organics

Sample No.	Type		Clienc	Matrix	Analysis	Due Date	
125611-001 125615-003 125616-003 125616-004 125628-007 125628-008			Burns & McDonnell Parsons Engineering Science, I Environmental Science & Engine Environmental Science & Engine Parsons Engineering Science, I Parsons Engineering Science, I	Water Water Water	TVH TVH/BTXE TVH TVH TVH/BTXE TVH/BTXE	22-MAY-96 22-MAY-96 23-MAY-96 23-MAY-96 23-MAY-96 23-MAY-96	
QC22221 QC22222 QC22223 QC22224 QC22225	LCS LCS MB MS MSD	of 125616-004 of 125616-004		Water Water Water Water			

Signed⁴

. Analyst: MMP Date: 5/20/96 Sequence Name: MAY 20 Page 63 Continued from Page Z File Prefix: 141 TKL Batch No.: 27767 Vial Comment Lims No. Wt/voi Std. File Stn. Sample Name STD Name NO. No No. See Premous 2 760 4 1855/my/ 39 1/vecd Paga 5 40 10:02 41 3 1 n. 102 Read and Understood by . Continued on Page:

Continued on Page: 65

Analyst: AMP /402 Date: 5/21/96 Sequence Name: MAY 2/ 64 Page File Prefix: 142 JKL Batch No.: Continued from Page: Wtvol Comment Std. Lims No. Vial File Stn. Sample Name NO. STD Name No. No 4.6 115012222 5mi 7055 11:22 GLW524CL A IS 450 mg/L 94W5Z432 B 1905 2000 M/L 1:5 BTXE 20 MY/1 4 96W52427 96W5Z123 IP-4 2080-4/L 941452036 Fax5 15 4 125.002228 2055 ーのクゴ 32 125600-001 36 · ni) 5

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Read and Understood by

Curtis & Tompkins, Ltd. Sample Batch Report

Ba 1 Number: 27666

Analysis : N/A Bgroup: : TVH Da Started: 18-MAY-96

Batched By : Malinda Ferguson

Department: Volatile Organics



Sample No. Ty	⁄pe		Client	Matrix	Analysis	Due Date	
125496-006			Montgomery Watson	Water	TVH	20-MAY-96	
125496-007			Montgomery Watson	Water	TVH	20-MAY-96	
125496-009			Montgomery Watson	Water	TVH	20-MAY-96	
125558-002			Subsurface Consultants	Water	TVH	20-MAY-96	
125572-001			Army Corps of Engineers, SPDL	Water	TVH	24-MAY-96	
125612-001			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-002			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-003			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-004			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-005			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-006			Burns & McDonnell	Water	TVH	20-MAY-96	
25612-007			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-008			Burns & McDonnell	Water	TVH	20-MAY-96	
25615-001			Parsons Engineering Science, I	Water	TVH/BTXE	22-MAY-96	
125615-002			Parsons Engineering Science, I	Water	TVH/BTXE	22-MAY-96	
125615-003			Parsons Engineering Science, I	Water	TVH/BTXE	22-MAY-96	
125615-004			Parsons Engineering Science, I	Water	TVH/BTXE	22-MAY-96	
25615-005			Parsons Engineering Science, I	Water	TVH/BTXE	22-MAY-96	
C22011 MB	1			Water			
C22012 LC	:s			Water			
C22013 LC				Water			
C22014 MS		125496-009		Water			
C22015 MS	D of	125496-009		Water			

Date: 5/20/96 Sequence Name: MAY 20 Page 62 Batch No.: 27707 File Prefix: 141 JKL Continued from Page: Sample Name File Stn. Wt/vol Comment Std. Lims No. Vial No No. NO. STD Name LCS.QCZ2170 16 PASSOF, 19/1/15/2388 CCV ISYSUMSIL PASS ... 96USZ432 B PASS / used 6A5 200043/L LCS & 22171 -1 8255/156 96US2206 B MB'GC 22172 ND BTXE 2015/L Retur Sonocite Lau! mss 1255m-009 41,WSZ4Z71B ms &c 22173 PASS BIXE 2019L 20 MSD AC 22174 10 125500-010 2.00 Overlange Run Merit 125465-001 TS450 15/L Clean-out System 10.5 % Over Range Run Mech 125472-001 18 1 Man out Jamples # # received Std #1) 112541,5-001 A1 Simple From #16+ received Std #6 10, 421 Posslused (Libration FOSS/rised on page SIA BK 01719. 60014 Rin Loa Book, lMeDH ... Clean Susien 25 MACH cleun system 1 125513-032 125549 -015 -C17-25524 -(0 -003 Tasilal 8:57 5/2/91

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5/21/92

Saich No. 2-7666 File Prefix 13 Continued from Page:	Analyst: CHL/CW Date: 5/20/96		196	Seque	nce Name: MA		Page_ 58 6-			
No. No 1	Batch No.:	: 2	-666		File Pr	efix: /39	C	Continued from Page: 1997		
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Batch N	10.: <u></u> 2	-7666	File Pre	fix: 139		ntinued from Page:	
File	311.	Sample Name	WUVOI	Comment		Lims No.	Vial
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Signed

Ravia Whalling Signed

SEQUENCE FILE: INSTRUMENT: BATCH:		·		MATRIX: I.C. DATE: ANALYSIS DATE	01/26	/96	
NTINUING CALIB	RATION VER AMOUNT (u	ıg/L)					
_	ACTUAL		CALC			STATUS	*
MTBE	20		31.58	157.9		FAIL	
BENZENE	20		18.8	94.0		PASS	
TOLUENE	20		19.98	99.9		PASS	
ETHYL BENZENE	20		19.84			PASS	
M, P-XYLENE	20		20.64			PASS PASS	
O-XYLENE	20		20.83	104.1		PASS	
TRIFLUOROTOLUENE	: 89	PASS		BROMOBENZENE:	89	PASS	
LABORATORY CONTR	OL SAMPLE: AMOUNT (u			FILE:142K001	QC222	22	
	ACTUAL		CALC	% REC		STATUS	*
MTBE	20		26.48			PASS	
BENZENE	20			90.4		PASS	
TOLUENE	20		19.01	95.1		PASS	
ETHYL BENZENE	20		18.68	93.4		PASS	
M, P-XYLENE	40		38.85				
O-XYLENE	20		19.91	99.6		PASS	
TRIFLUOROTOLUENE	: 91	PASS		BROMOBENZENE:	92	PASS	
CONTINUING CALIB	RATION VER	RIFICAT	TION:	FILE:142K012			
	AMOUNT (u						
	ACTUAL	(CALC	% REC		STATUS	*
MTBE	20		33.8			FAIL	
BENZENE	20		16.91			PASS	
TOLUENE	20		19.19			PASS	
ETHYL BENZENE	20		17.86				
M, P-XYLENE	20		19.79			PASS	
O-XYLENE	-20		21.05	105.3		PASS	
TRIFLUOROTOLUENE	: 93	PASS		BROMOBENZENE:	92	PASS	
RECOVERY CCV LIMITS LCS CURTIS & TOMPKIN	80-120%	(MTBE (MTBE	80-120%) 65-135%)	TRIFLUOROTOI BROMOBENZENI	LUENE E (62-	(58-130 131%))%)

SEQUENCE FILE: INSTRUMENT:	MAY21 GC04	MATRIX: WATE	
	27719	I.C. DATE: 01/2 DATE OF ANALYSIS:	•
CONTINUING CALIB	RATION VERIFICATION: AMOUNT (ug/Kg)	· ·	
1,2,3-TMB 1,2,4-TMB 1,3,5-TMB	ACTUAL CALC 20.0 20 20.0 19 20.0 19	.8 98.8	PASS
TRIFLUOROTOLUENE	: 91 PASS	BROMOBENZENE: 92	PASS
CONTINUING CALIB	RATION VERIFICATION:	FILE: G:\GC04\142k	(012
	AMOUNT (ug/Kg)		
1,2,3-TMB 1,2,4-TMB 1,3,5-TMB	AMOUNT (ug/Kg) ACTUAL CALC 20.0 21	.5 112.3	STATUS *
1,2,4-TMB	AMOUNT (ug/Kg) ACTUAL CALC 20.0 21 20.0 22 20.0 22	.9 109.5 .5 112.3	STATUS * PASS PASS PASS

						CC	Curtis & Tompkins, Ltd.
		ACTUAL	(CALC	% REC		STATUS *
	MTBE	20		38.2	191.0		FAIL
	BENZENE	20		18.4	92.0		PASS
	TOLUENE	20		19.1	95.5		PASS
1	ETHYL BENZENE	20		19.2	96.0		PASS
	M, P-XYLENE	20		19.7	98.5		PASS
	O-XYLENE	20		20.1	100.5		PASS
	TRIFLUOROTOLUENE	93	PASS		BROMOBENZENE:	96	PASS
	RECOVERY CCV LIMITS LCS CURTIS & TOMPKINS	85-115% 80-120% S, LTD	•	80-120%) 65-135%)			•

Curtis & Tompkins, Ltd

MATRIX: SEQUENCE FILE: MAY18 WATER INSTRUMENT: GC04 I.C. DATE: 01/26/96

BATCH: 27666 DATE OF ANALYSIS: 05/18/96

CONTINUING CALIBRATION VERIFICATION: FILE: G:\GC04\139K003

AMOUNT (ug/Kg)

	ACTUAL	CALC	% REC	STATUS :
1,2,3-TMB	20.0	18.6	93.0	PASS
1,2,4-TMB	20.0	18.8	93.9	PASS
1,3,5-TMB	20.0	19.1	95.3	PASS

TRIFLUOROTOLUENE: 94 PASS BROMOBENZENE: 95 PASS

CONTINUING CALIBRATION VERIFICATION: FILE: G:\GC04\139K015

AMOUNT (ug/Kg)

	ACTUAL	CALC	* REC	STATUS *
1,2,3-TMB	20.0	18.4	92.0	PASS
1,2,4-TMB	20.0	19.0	94.9	PASS
1,3,5-TMB	20.0	20.2	100.9	PASS

TRIFLUOROTOLUENE: 95 PASS BROMOBENZENE: 97 PASS

CONTINUING CALIBRATION VERIFICATION: FILE: G:\GC04\139K032

AMOUNT (ug/Kg)

	ACTUAL	CALC	% REC	STATUS *
1,2,3-TMB	20.0	19.3	96.5	PASS
1,2,4-TMB	20.0	19.8	99.2	PASS
1,3,5-TMB	20.0	20.9	104.6	PASS

TRIFLUOROTOLUENE: 94 PASS BROMOBENZENE: 96 PASS

CONTINUING CALIBRATION VERIFICATION: FILE: G:\GC04\139K040

AMOUNT (ug/Kg)

	ACTUAL	CALC	% REC	STATUS *
1,2,3-TMB	20.0	20.0	100.2	PASS
1,2,4-TMB	20.0	20.2	101.0	PASS
1,3,5-TMB	20.0	21.0	105.0	PASS

TRIFLUOROTOLUENE: 93 PASS BROMOBENZENE: 96 PASS

RECOVERY CCV 85-115% TRIFLUOROTOLUENE (57-131%)

LIMITS 1,2,3-TMB 80-120% BROMOBENZENE (61-132%)

CURTIS & TOMPKINS, LTD HCR Group Name :
Calibrating Area versus Amount Using a 1st Order Fit
Amounts Will Not Be Scaled Prior To The Regression
Weighting Factor For the Regression: 1
Calibration Levels:

Level Name	_ Amount _	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2,5000	0.00	0.00			0
2	12.5000	22065.80	4676.22			1
3	50.0000	49944.10	10528.00			1
4	100.0000	101038.30	21488.12			1
5	500.0000	505039.59	104463.09		•••••	1
6	750.0000	815365.13	168480.52	• • • • • • • • • • • • • • • • • • • •		1
7	1500.0000	1485334.00	303323.59			1

Calibration Curve : $y = (8397.2979) + (1001.7173)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99785

1,3,5-TMB

Retention Time : 8.280 min Search Window: 2 sec, 2 % Reference Component: TRIFLUOROTOLUENE (Find Largest Peak)

Group Name : Calibrating Area wergus Ame

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1 Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2.5000	0.00	0.00			0
2	12.5000	18122.60	3942.51			1
3	50.0000	42761.40	9114.75			1
4	100.0000	91438.99	19152.92	•••••		1
5	500.0000	454541.84	93784.82			1
6	750.0000	726311.50	149146.02			1
7	1500.0000	1360324.63	275414.16			1

Calibration Curve : $y = (3913.8481) + (914.6154)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99890

Curtis & Tompkins, Ltd

MATRIX: SEQUENCE FILE: MAY18 WATER I.C. DATE: 01/26/96 GC04 INSTRUMENT: ANALYSIS DATE: 05/18/96 BATCH: 27666 FILE:139K003 CONTINUING CALIBRATION VERIFICATION: AMOUNT (uq/L) % REC STATUS * CALC ACTUAL 36.9 184.5 MTBE 20 FAIL PASS 17.9 89.5 BENZENE 20 PASS 92.5 20 18.5 TOLUENE 20 18.4 92.0 PASS ETHYL BENZENE 94.5 PASS 20 18.9 M,P-XYLENE 96.0 PASS 20 19.2 O-XYLENE TRIFLUOROTOLUENE: 94 PASS BROMOBENZENE: 95 PASS _____ _____ FILE:139K004 QC22013 LABORATORY CONTROL SAMPLE: AMOUNT (ug/L) ACTUAL CALC % REC STATUS * 168.5 33.7 FAIL 20 MTBE 100.0 PASS 20 20 BENZENE 101.5 20.3 PASS 20 TOLUENE PASS 20 19.8 99.0 ETHYL BENZENE 40 40.7 101.8 PASS M, P-XYLENE 20.6 103.0 PASS O-XYLENE 20 BROMOBENZENE: 94 PASS TRIFLUOROTOLUENE: 95 PASS ______ CONTINUING CALIBRATION VERIFICATION: FILE: 139K015 AMOUNT (ug/L) CALC STATUS *
FAIL
PASS % REC 172.5 ACTUAL 34.5 MTBE 20 17.6 88.0 BENZENE 20 92.5 PASS 18.5 TOLUENE 20 20 ETHYL BENZENE 17.9 89.5 PASS 18.7 93.5 M, P-XYLENE 20 94.5 20 18.9 O-XYLENE TRIFLUOROTOLUENE: 95 PASS BROMOBENZENE: 97 PASS CONTINUING CALIBRATION VERIFICATION: FILE:139K032 AMOUNT (ug/L) STATUS * CALC % REC ACTUAL 41.9 209.5 MTBE 20 96.0 PASS 20 19.2 BENZENE 19.9 99.5 PASS 20 TOLUENE ETHYL BENZENE 20 19.6 98.0 PASS 20 20.3 101.5 PASS M, P-XYLENE 20.7 O-XYLENE 20 103.5 PASS TRIFLUOROTOLUENE: 94 PASS BROMOBENZENE: 96 PASS _____ _____

CONTINUING CALIBRATION VERIFICATION: FILE:139K040 AMOUNT (ug/L)

Level Name	Arount	Area	aeignt	ישים אבשף.	.SIU AME.	≆ keptica:∟,
1	2.5000	1934.40	430.00			1
2	12.5000	12493.00	2831.61			1
3	50.0000	43688.30	9561.35		*********	1
4	100.0000	100247.80	21566.70			1
5	500.0000	472436.91	99938.37			1
6	750.0000	702088.00	149475.34	•••••		1
7	1500,0000	1459776.38	309126.72			1



Calibration Curve : $y = (-407.5063) + (959.0111)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99951

BROMOBENZENE

Retention Time : 6.785 min Search Window: 5 sec, 5 % Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Group Name

Calibrating Height versus Amount Using a Pt. to Pt. Fit Amounts Will Not Be Scaled Prior To The Regression

Weighting Factor For the Regression: 1

Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	 # Replicates
8	2.5000		167400.42		 1

Sample File: G:\GC04\CONFIRM\CTMB.smp
Created by:
on: 5/12/89 2:54 PM
Edited by:
on: 5/20/96 2:37 PM
Number Of Times Edited: 0

Calibration Dute: "4/

Sample Description

Default Injection Volume = 1.0000 ul

An External Standard Calibration Will Be Used

Unknown Peaks Will Be Quantitated Using A Response Factor of 1000000.0000

Component Information:

TRIFLUOROTOLUENE

Retention Time : 2.842 min Search Window: 5 sec, 5 % Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Group Name :

Calibrating Height versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression

Weighting Factor For the Regression: 1

Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	
8	2.5000	370592.59	78089.03		 1

Calibration Curve : $y = (0.0000) + (0.0000)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.00000

BROMOBENZENE

Retention Time : 6.801 min Search Window: 5 sec, 5 % Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Group Name

Calibrating Height versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1 Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	# Replicates
8	2.5000	1212933.88			 1

Calibration Curve : $y = (0.0000) + (0.0000)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.00000

1,2,3-TMB

Retention Time : 7.290 min Search Window: 2 sec, 2 % Reference Component: TRIFLUOROTOLUENE (Find Largest Peak)

Group Name :

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1

Calibration Levels:

. ucioni il						
Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2.5000	7595.00	945.18			1
2	12.5000	32122.20	6113.12			1
3	50.0000	69130.00	13790.98		• • • • • • • • • • • • • • • • • • • •	i
4	100.0000	139994.77	28033.53			i
5	500.0000	647813.19	131498.53			i
6	750.0000	1015770.81	210788.63			i
7	1500.0000	1867216.88	372549.75			i

Calibration Curve : $y = (18542.1094) + (1252.0919)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99842

1,2,4-TMB

Retention Time : 7.750 min Search Window: 2 sec, 2 % Reference Component: TRIFLUOROTOLUENE (Find Largest Peak)

044

Sample File : G:\GC04\CONFIRM\CBTXE3.smp

on: 5/12/892:54 PM Created by : on: 1/26/96 9:10 AM Edited by

Number Of Times Edited: 23

range = 10

e Description fault Injection Volume = 1.0000 ul

An External Standard Calibration Will Be Used

Unknown Peaks Will Be Quantitated Using A Response Factor of 1000000.0000

Component Information :

MTBE

Search Window: 5 sec, 5 % Retention Time : 1.154 min Reference Component: TRIFLUOROTOLUENE (Find Largest Peak)

Group Name

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1

Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
2	12.5000	2414.90	644.42			1
3	50,0000	9064.40	2289.76		• • • • • • • • • • • • • • • • • • • •	1
4	100.0000	23609.60	5872.21		• • • • • • • • • • • • • • • • • • • •	1
5	500.0000	116073.31	28264.73			, 1
6	750,0000	164030.30	39891.57			1
7	1500.0000	366475.81	89698.11			1

 $y = (-4055.2905) + (242.4035)x + (0.0000)x^2 + (0.0000)x^3$ Calibration Curve Correlation Coefficient: 0.99733

BENZENE

Retention Time : 2.064 min Search Window: 5 sec, 5 % Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1/x

Calibration Levels:

ac-011 -0						
Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2.5000	2895.40	784.98			1
2	12.5000	18379.90	7,23.00			1
3	50.0000	60970.80	13712.04			1
4	100.9000	135749.00	22,000			1
5	500.0000	611710.63	155399.27			1
6	750.0000	890909.00	227964.16			1
7	1500.0000	1804479.00	420202.00			ı

 $y = (713.3722) + (1207.5815)x + (0.0000)x^2 + (0.0000)x^3$ Calibration Curve Correlation Coefficient: 0.99979

TRIFLUOROTOLUENE

Search Window: 5 sec, 5 % : 2.824 min Retention Time Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Calibrating Height versus Amount Using a Pt. to Pt. Fit Amounts Will Not Be Scaled Prior To The Regression

Weighting Factor For the Regression: 1

Calibration Levels:

Level Name	Amount	Area	Height		# Replicates
8	2.5000				

Search Window: 5 Sec, 5 6 Retention Time : 3.718 min Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Calibrating Area versus Amount Using a 1st Order Fit

Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1/x

Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2.5000	2721.80	673.13			1
2	12.5000	15965.00	3825.36	•••••		1
3	50.0000	53769.50	12606.48			1
4	100.0000	120509.40	27888.65			1
5	500.0000	558034.13				1
6	750.0000	828766.44	188109.39			1
7	1500.0000	1682478.50	380409.78	*********	••••	•

: $y = (312.3202) + (1118.3734)x + (0.0000)x^2 + (0.0000)x^3$ Calibration Curve Correlation Coefficient: 0.99991

ETHYLBENZENE

Search Window: 5 sec, 5 % Retention Time : 5.378 min Reference Component: BROMOBENZENE (Find Closest Peak)

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1/x

Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2.5000	1964.51	447.51	•••••	••••••	1
2	12.5000	12842.49	2931.25		•••••	1
3	50.0000	44289.00	9898.41			1
4	100.0000	100726.29	22130.27		•••••	1
5	500.0000	468847.34	101668.99			1
6	750.0000	710752.25	154158.20			1
7	1500,0000	1446143.63	312055.94			1

Calibration Curve : $y = (-268.4567) + (956.2418)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99982

m,p-XYLENE

Retention Time : 5.526 min Search Window: 5 sec, 5 % Reference Component: BROMOBENZENE (Find Closest Peak)

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1/x

Calibration Levels:

ation ne	V=13.					
Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2,5000	2762.99	636.82	•••••	•••••	1
2	12,5000	16055.68	3590.60		•••••	1
3	50.0000	55524.80	11979.87		***********	1
4	100.0000	126116.27	26609.83			1
5	500.0000	583491.13	121608.00			1
6	750.0000	869212.44	183989.94			1
7	1500.0000	1765456.25	374175.31			1

Calibration Curve : $y = (110.2872) + (1172.5034)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99990

o-XYLENE

Retention Time : 5.998 min Search Window: 5 sec, 5 %

Reference Component: BROMOBENZENE (Find Closest Peak)

Group Name

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1/x antibuncian tarrata.

Sample File : G:\GC04\TMB.smp on: 5/12/892:54 PM Created by : Edited by : on: 5/20/96 2:37 PM Number Of Times Edited: 0 Culibration Date: 1/26/96 CW 5/20/96

Sample Description

Default Injection Volume = 1.0000 ul

An External Standard Calibration Will Be Used

Unknown Peaks Will Be Quantitated Using A Response Factor of 1000000.0000

Component Information:

TRIFLUOROTOLUENE

Retention Time : 2.413 min Search Window: 5 sec, 5 % Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Group Name

Calibrating Height versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1

Calibration Levels:

Level Name ISTD Resp. ISTD Amt. # Replicates Amount Height Area R 2.5000 344106.19 76849.11 ----

Calibration Curve : $y = (0.0000) + (0.0000)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.00000

BROMOBENZENE

Retention Time : 6.162 min Search Window: 5 sec, 5 % Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Calibrating Height versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1

Calibration Levels:

Level Name Amount ISTD Resp. Area Height ISTD Amt. # Replicates R 2.5000 1152127.38 239019.05 ------

Calibration Curve : $y = (0.0000) + (0.0000)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.00000

1,2,3-TMB

Retention Time : 6.930 min Search Window: 2 sec, 2 % Reference Component: TRIFLUOROTOLUENE (Find Largest Peak)

Group Name

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1

Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2.5000	4676.70	856.68			1
2	12.5000	34262.44	6442.38		•••••	1
3	50.0000	69879.58	13413.39			1
4	100.0000	133746.56	26691.03	•••••		1
5	500.0000	599672.38	124397.88			1
6	750,0000	973281.25	203010.16			1
7	1500.0000	1709913.00	355363.03			1

Calibration Curve $y = (23975.0020) + (1151.8377)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99608

2.4-TMB

Retention Time : 7.360 min Search Window: 2 sec, 2 % Reference Component: TRIFLUOROTOLUENE (Find Largest Peak)

039

Group Name :
Calibrating Area versus Amount Using a 1st Order Fit
Amounts Will Not Be Scaled Prior To The Regression
Weighting Factor For the Regression: 1

Calibration Levels:

Level Name	. Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2,5000	3044.20	607.84			1
2	12.5000	24689.41	4804.05			1
3	50.0000	49810.80	10129.09			1
4	100.0000	100136.74	20445.47			1
5	500.0000	461291.56	96790.13			1
6	750,0000	744361.63	157172.66			1
7	1500.0000	1326502.38	279349.44			1

Calibration Curve : $y = (15042.8486) + (893.4946)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99688

1,3,5-TMB

Retention Time : 7.830 min Search Window: 2 sec, 2 % Reference Component: TRIFLUOROTOLUENE (Find Largest Peak)

Group Name

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1

Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2.5000	3732.67	701.88			1
2	12.5000	24769.70	4488.30			1
3	50.0000	51657.16	9212.12			1
4	100,0000	94843.96	18542.52			1
5	500.0000	424592.44	87453.02			1
6	750.0000	672444.44	139600.66			1
7	1500.0000	1215791.50	254956.55			1

Calibration Curve : $y = (16068.7236) + (814.8716)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99774 Retention Time : 3.369 min Search Window: 5 sec, 5 % Reference Component: TRIFLUOROTOLUENE (Find Closest Peak) Group Name : Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1/x



Ca:	libration Le	vels:	Ares	Height	ISTD Resp.	ISTD Amt.	# Replicates
	1	2.5000	4033.10	975.29			1
,	2	12.5000	18550.40	4424.93	•••••		1
	3	50.0000	56661.80	13289.72		•••••	1
	4	100.0000	122639.10	28695.62	•••••		1
	5	500.0000	543820.63	128552.16	•••••	•••••	1
	. 6	750.0000	806585.88	192251.23		•••••	1
	7	1500.0000	1652944.88	393321.59		•••••	1

Calibration Curve : y = (2061.5588) + (1094.6157)x + (0.0000)x² + (0.0000)x³ Correlation Coefficient: 0.99977

ETHYLBENZENE

Retention Time : 5.015 min Search Window: 5 sec, 5 % Reference Component: BROMOBENZENE (Find Closest Peak)

Group Name :

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1/x

Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2.5000	2966.51	634.66			1
ż	12.5000	15320.36	3373.62			1
3	50.0000	47364.98	10348.65		•••••	1
4	100.0000	102356.15	22432.36			1
5	500.0000	460443.63	101715.31			1
6	750.0000	696296.38	154284.42			1
7	1500.0000	1417643.38	314905.75			1

Calibration Curve : $y = (1190.6255) + (937.9269)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99981

m, p-XYLENE

Retention Time : 5.169 min Search Window: 5 sec, 5 % Reference Component: BROMOBENZENE (Find Closest Peak)

Group Name

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1/x

Calibration Levels:

ation Le	vers:					
Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2.5000	4333.96	915.03			1
2	12.5000	19232.25	4096.34			1
·3	50.0000	58617.82	12277.18			1
4	100.0000	126606.71	26623.63			1
5	500.0000	563235.31	120459.57	•••••		1
6	750.0000	836746.75	182632.80			1
7	1500.0000	1700683.75	3/4/62.63		**********	1

Calibration Curve : $y = (2291.4168) + (1129.8170)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99986

o-XYLENE

Retention Time : 5.578 min Search Window: 5 sec, 5 % Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Group Name

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1/x

037

Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2,5000	2662.90	611.05	•••••		1
1	12,5000	14951.30	3187.77		•••••	1
7	50.0000	46354.30	9738.74		•••••	1
4	100,0000	100849.13	21364.83	•••••		1
5	500.0000	455173.00	97308.00		•••••	1
6	750.0000	670957.81	145525.41	•••••]
7	1500.0000	1383192.13	300450.78	••••••		1

: $y = (1010.3149) + (914.9463)x + (0.0000)x^2 + (0.0000)x^3$ Calibration Curve Correlation Coefficient: 0.99971

BROMOBENZENE

Search Window: 5 sec, 5 % : 6.156 min Retention Time

Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Group Name

Calibrating Height versus Amount Using a Pt. to Pt. Fit

Amounts Will Not Be Scaled Prior To The Regression

Weighting Factor For the Regression: 1

Calibration Levels:

Level Name	Amount	Area			# Replicates
8	2.5000	787998.31	166316.73	•••••	 1 .



Lab #: 125615

BATCH QC REPORT

Page 1 of 1

BTXE

Parsons Engineering Science, Inc. Client:

Analysis Method: EPA 8020

Project#: 722450.14020

Prep Method:

EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Prep Date:

05/21/96

Water Matrix: Batch#: 27719 ug/L Units:

Analysis Date:

05/21/96

Diln Fac: 1

LCS Lab ID: QC22222

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	18.08	20	90	80-120
Toluene	19.01	20	95	80-120
Ethylbenzene	18.68	20	93	80-120
m,p-Xylenes	38.85	40	97	80-120
o-Xylene	19.91	20	100	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	91	58-130		
Bromobenzene	92	62-131		

Column to be used to flag recovery and RPD values with an asterisk

Values outside of QC limits

ike Recovery: 0 out of 5 outside limits

Sample File : G:\GC04\BTXE3.smp

Created by : on : 5/12/89 2:54 PM Edited by : on : 1/26/96 9:09 AM

Number Of Times Edited: 63

range = 2

Sample Description :

Default Injection Volume = 1.0000 ul

An External Standard Calibration Will Be Used

Unknown Peaks Will Be Quantitated Using A Response Factor of 1000000.000

intensity = 1

. Component Information :

MTBE

Retention Time : 1.114 min Search Window: 5 sec, 5 % Reference Component: TRIFLUOROTOLUENE (Find Largest Peak)

Group Name

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1

Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
2	12.5000	4020.66	989.61		****	1
3	50.0000	11330.50	2794.52			1
4	100.4000	25637.00	6483.12	•••••		1
5	500.0000	116573.10	28895.31			1
6	750.0000	160983.00	40867.50			1
7	1500.0000	356797.59	91765.27			1

Calibration Curve : $y = (-1295.3915) + (234.5134)x + (0.0000)x^2 + (0.0000)x^3$ Correlation Coefficient: 0.99734

BENZENE

Retention Time : 1.815 min Search Window: 5 sec, 5 % Reference Component: TRIFLUCROTOLUENE (Find Closest Peak)

Group Name

Calibrating Area versus Amount Using a 1st Order Fit Amounts Will Not Be Scaled Prior To The Regression Weighting Factor For the Regression: 1/x

Calibration Levels:

. acioni ne						
Level Name	Amount	Area	Height	ISTD Resp.	ISTD Amt.	# Replicates
1	2.5000	4039.30	1226.65			1
2	12.5000	20863.00	6227.44			1
3	50.0000	62341.00	18491.57			1
4	100.0000	135119.70	40068.50			1
5	500.0000	596198.13	178782.27		• • • • • • • • • • • • • • • • • • • •	1
6	750.0000	882501.81	265308.38			1
7	1500,0000	1801279.88	541700.50			1

- Calibration Curve : y = (2096.3135) + (1196.4558)x + (0.0000)x² + (0.0000)x³ Correlation Coefficient: 0.99981

TRIFLUOROTOLUENE

Retention Time : 2.411 min Search Window: 5 sec, 5 % Reference Component: TRIFLUOROTOLUENE (Find Closest Peak)

Group Name

Calibrating Height versus Amount Using a Pt. to Pt. Fit Amounts Will Not Be Scaled Prior To The Regression

Weighting Factor For the Regression: 1

Calibration Levels:

Level Name	Amount	Area	Height	ISTD Resp.	# Replicates
8	2.5000		66228.97		1



ab #: 125615

BATCH QC REPORT

Page 1 of 1

BTXE

lient: Parsons Engineering Science, Inc. Analysis Method: EPA 8020

Project#: 722450.14020 Location: Beale AFB

Prep Method:

EPA 5030

METHOD BLANK

Matrix: Water Prep Date:

05/21/96

Batch#: 27719 Analysis Date:

05/21/96

Units: ug/L Diln Fac: 1

MB Lab ID: QC22223

Analyte	Result	A.
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene	88	58-130
romobenzene	87	62-131

Page 1 of 1

BATCH QC REPORT

Lab #: 125615

BTXE

Client: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

Project#: 722450.14020

Water

Prep Method: EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Prep Date:

Prep Date: 05/18/96 Analysis Date: 05/18/96

Batch#: 27666 Units: ug/L Diln Fac: 1

Matrix:

LCS Lab ID: QC22013

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	20	20	100	80-120
Toluene	20.3	20	102	80-120
Ethylbenzene	19.8	20	99	80-120
m,p-Xylenes	40.7	40	102	80-120
o-Xylene	20.6	20	103	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	95	58-130	· · · · · · · · · · · · · · · · · · ·	
Bromobenzene	94	62-131		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 5 outside limits



BTXE

client: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

Project#: 722450.14020 Location: Beale AFB

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
125615-005 TRIP BLANK	27666	05/15/96	05/19/96	05/19/96	

Matrix: Water

Analyte	Units	125615-005	
Diln Fac:		1	
Benzene	ug/L	<0.5	
Toluene	ug/L	<0.5	
Ethylbenzene	ug/L	<0.5	
m,p-Xylenes	ug/L	<0.5	
o-Xylene	ug/L	<0.5	
1,2,3-Trimethylbenzene	∍ ug/L	<0.5	
2,4-Trimethylbenzene	e ug/L	<0.5	
3,5-Trimethylbenzene	e ug/L	<0.5	
Surrogate			
Trifluorotoluene	%REC	95	
Bromobenzene	%REC	97	



Lab #: 125615

BATCH QC REPORT

Page 1 of 1

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B	Τ.	А	Ľ

Client: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

Project#: 722450.14020 Location: Beale AFB

Prep Method:

EPA 5030

METHOD BLANK

Matrix: Water

Prep Date:

05/18/96

Batch#: 27666 ug/L

Analysis Date: 05/18/96

Units: Diln Fac: 1

MB Lab ID: QC22011

Analyte	Result	
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	

Surrogate	%Rec	Recovery Limits
Trifluorotoluene	96	58-130
Bromobenzene	95	62-131

BTXE Data

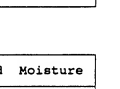


BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB Analysis Method: EPA 8020

Prep Method: EPA 5030



Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
125615-001 BE-MP10-01	27666	05/14/96	05/19/96	05/19/96	
125615-002 BE-MP12-01	27666	05/15/96	05/19/96	05/19/96	
125615-003 BE-MP13-01	27719	05/14/96	05/21/96	05/21/96	
125615-004 BE-MP47-01	27666	05/14/96	05/19/96	05/19/96	

Matrix: Water

Analyte	Units	125615-001	125615-002	125615-003	125615-004
Diln Fac:		1	1	5	1
Benzene	ug/L	<0.5	<0.5	<2.5	<0.5
Toluene	ug/L	<0.5	<0.5	9.2	<0.5
Ethylbenzene	ug/L	20	<0.5	22	<0.5
m,p-Xylenes	ug/L	19	<0.5	35	<0.5
o-Xylene	ug/L	2.9	<0.5	4.9	<0.5
1,2,3-Trimethylbenzen	e ug/L	<0.5	<0.5	, 120	<0.5
1,2,4-Trimethylbenzen	e ug/L	3.6	<0.5	<2.5	<0.5
1,3,5-Trimethylbenzen	e ug/L	7.8	<0.5	59	<0.5
Surrogate					
Trifluorotoluene	%REC	113	93	119	95
Bromobenzene	%REC	106	94	91	98

Analyst: <u>AMP/HCR</u> Date: <u>5/21/96</u> Sequence Name: <u>MAV 2/</u> Page 65 Batch No.: 1 27723 142 JKL File Prefix:_ Continued from Page: 64 File Stn. Sample Name Wt/vol Comment Lims No. No No. NO. STD Name 125519-006 40 PASSINSED 44 8:11 Spotox All tuns received std # Continued on Page:__ Read and Understood by

Curtis & Tompkins, Ltd. Sample Batch Report

Batch Number: 27719 Date Started: 21-MAY-96 Batched By : Anna Maria Pajarillo

Analysis : N/A Bgroup: : TVH Department: Volatile Organics

Sample No.	Туре		Client	Matrix	Analysis	Due Date	
125611-001			Burns & McDonnell	Water	TVH	22-MAY-96	
125615-003			Parsons Engineering Science, I	Water	TVH/BTXE	22-MAY-96	
125616-003			Environmental Science & Engine	Water	TVH	23-MAY-96	
125616-004			Environmental Science & Engine	Water	TVH	23-MAY-96	
125628-007			Parsons Engineering Science, I	Water	TVH/BTXE	23-MAY-96	
125628-008			Parsons Engineering Science, I	Water	TVH/BTXE	23-MAY-96	
QC22221	LCS		•	Water			
QC22222	LCS			Water			
QC22223	MB			Water			
QC22224	MS	of 125616-004		Water			
QC22225	MSD	of 125616-004		Water			

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No. No. STD. Name 39			· · · · · · · · · · · · · · · · · · ·	7		I .		
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Analyst	: AMP	HCZ Date: 5/21/96	Sequenc	ce Name: <u>MAY 2 /</u>		Pag	ge	64
Batch N	ار م	7719/27723	File Pre	fix: 142 IKL	Cor	tinued from P	age:	
File	Stn.	Sample Name	Wt/vol	Comment	Std.	Lims No.		Vial
No.	No				NO.	STD Nar	ne	
1	4,6	LC5 AC22222	5mi	7055 11:22	1 /	94.11524		A
2		MB. D(22223	1	<u> </u>		IS 450 m		
3		125616-004	1/1		2	94 W5 24	<u> 32</u>	ゟ
. 4		MS, B(22224				605 2000 m		
5		M5D A(22225	I V		3	96W522	06	B
6		125615-003	1 1	1:5		BTXE 20 M	14/1	
7		125611-001		1:2	4	9645242	7	
8		125628-007				BIXE 20 Mg	1/2 2	2.1
9		125429-008	<i>V</i>		5	96W5212	3	
10	1	12 5614 -003				JP-4 2000	29/2	
11	2	CCV	₩		6	96115203	6	
12	3,6	CCU	101	Pass		THE 20 MI	2	
/3	2	LCS . AC 22237	1	7a35				
14	3	CCV	/	<u> </u>				
15	4	LC5.0(22238		Dass				
16	15	!CCV		Pass				
17		MB 18(22239						
18		125600-002				•		
19	7	MS QLZ2240						
20	4	MSD Q(22241					1	
21		125524-023						
22		1 - 024						
23		-025						
27		1 -026	<u> </u>					
25		V -027						
26		-028						
27	2	CCV						
23	3	CCV						
29	5	CCV	<u> </u>					
30	2	CrV		Fass				
3/	3	CCV		Pass				
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Read and Understood by

5/20/01

Curtis & Tompkins, Ltd. Sample Batch Report

Batch Number: 27666

Started: 18-MAY-96

B ... ched By : Malinda Ferguson

Analysis : N/A Bgroup: : TVH

Department: Volatile Organics

V	

Sample No.	Туре		Client	Matrix	Analysis	Due Date	
125496-006			Montgomery Watson	Water	TVH	20-MAY-96	
125496-007			Montgomery Watson	Water	TVH	20-MAY-96	
125496-009			Montgomery Watson	Water	TVH	20-MAY-96	
125558-002			Subsurface Consultants	Water	TVH	20-MAY-96	
125572-001			Army Corps of Engineers, SPDL	Water	TVH	24-MAY-96	
125612-001			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-002			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-003			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-004			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-005			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-006			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-007			Burns & McDonnell	Water	TVH	20-MAY-96	
125612-008			Burns & McDonnell	Water	TVH	20-MAY-96	
125615-001			Parsons Engineering Science, I	Water	TVH/BTXE	22-MAY-96	
125615-002			Parsons Engineering Science, I		TVH/BTXE	22-MAY-96	
125615-003			Parsons Engineering Science, I		TVH/BTXE	22-MAY-96	
125615-004			Parsons Engineering Science, I		TVH/BTXE	22-MAY-96	
125615-005			Parsons Engineering Science, I	Water	TVH/BTXE	22-MAY-96	
QC22011	MB			Water			
QC22012	LCS			Water			
QC22013	LCS			Water			
QC22014	MS	of 125496-009		Water			
QC22015	MSD	of 125496-009		Water			

KM126

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Date: 5/20/96 Sequence Name: MAY 20 Page 62 Batch No.: 27707 File Prefix: 141 JKL Continued from Page: File Stn. Sample Name Wt/voi Comment Std. Lims No. Vial No. No NO. STD Name 2 LCS, &CZ2170 119 961152388 13 CCV 15450MSIL PASS . 96WSZY32 B PASS / used GAS 2000 MS/L LCS AC 22171 I Passivsed 96US2206 B MB'BC 22172 ND BTXE 2015/L mss 125500-009 Relun Sowocate Law 91.WSZ4Z718 ms &c 22173 PASS BIXE 2015/L 20 MSD GC 22174 96WSZ123 10 125500-010 JP-4 2,00 M5/L 1 -009 6 96WS2466 125465 -001 Overlange Run Mech IS450 45/L 1B Clean-out System 10.5 % 101 en Rance Run Medt 125472-001 1 13 19 1 regnat Danjoles #7-15 15 #1-15 system received Sta #(1) 12541,5-001 1:25 Mech O.R. 17 P11 Sumples from #16+ 13 19 necented Std #6 19 CCV In that Calibration on page 51 de Pass/used BK 17219 Run LOABOOK 125465-001 IMeDH ... clean system 1125472-1001 1:25 Mech Cleun system 125513-032 125549 -015 -617₋ -CP7 125513-033 125524 -CC1 $-\infty Z$ -003 115 EEV 8:37 5/2/01 Continued on Page: 63 Read and Undepetood by

Analyst: CHR/CW Date: 5/20/96 Sequence Name: MAY 18 Page: 58: -Batch No.: 27666 File Prefix: 139 Continued from Page: * ** Lims No. File Sample Name Wt/vol Stn. Comment Std. 5/18/96 1210 NO. STD No. No Name ' Ł 2 CCU, 0C22012 pass/usel 96452388 C 57000 IS 450 mg/L 36 CCU, BX+ THE'S pun/usul 96WS2310-13 LCS QC 22013 Gas 2000 m/ MR QC 22011 26 BOLLSWAP 125496-9 BTXIE 20m/L MSS passad MS QC 22014 96052427 A us) ac 22015 BTXF 20m/6 Son 125572-1 95WS/57 0 9 stoddard 2000m/L 125612-2 u 96452036 L -5 13 oussed/used CCU 14 ccv 15 ccu CeU 36 12 IB 18 125612-6 19 20 21 125496-6 上 - 2 23 24 125615 - 2 1 -3 25 طله 27 28 29 cev 30 3.6 ccil حذلا 31 32 CCU pigsed/used IB 33 34 125612-1 1/50 125555-2 35 1/02,622 L 36 nossel/went SW 2 CCU 37 3.6 CCV Sul Read and Understood by

Continued on Page: 57

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Signed

Analyst: CHR / CW Date: 5/20/76 Sequence Name: MACIS Page 59 Batch No .: 27666 File Prefix: 139 Continued from Page: 58 Lims No. اندارد اندارد اندارد File Sample Name Wt/vol Vial Comment STA No. NO. STD Name 39 520 cev See previous 40 CCU 41 Jed 5/19/86 STOOD 0829 STO #1 BK 07-19 51 Continued on Page:____ Read and Understood by

Signed

5/20/86

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Date

CURTIS & TOMPKINS, LTD

BROMOBENZENE (BB): 70-122 %

INSTRUMENT: GC04 DATE OF INJECTION21-MAY-96

SEOUENCE: MAY20-21 MATRIX: WATER I.C.DATE: 01/25/96 BATCH: 27719

RESULT FILE: G:\GC04\141J038 LABORATORY CONTROL SAMPLE:

QC22221

AMOUNT (ug/L) ACTUAL CALC % REC STATUS

2000 90.1% IN COMPLIANCE 1802

SURROGATES, %REC : TFT = 92 PASS BB = 94 PASS

CONTINUING CALIBRATION CHECK: RESULT FILE: G:\GC04\142J013

AMOUNT (ug/L) ACTUAL CALC % REC STATUS

2000 1926.99 96.3% IN COMPLIANCE LUFT

SURROGATES, %REC : TFT = 95 PASS BB = 95 PASS

MSS sample: 125616-004 FILE 142J003

LUFT (ug/L)

SURROGATES, %REC : TFT = 84 PASS BB = 86 PASS -----

MATRIX SPIKE: ACTUAL CALC % REC STATUS 142J004 LUFT (ug/L) 2000 1796.42 89.8% IN COMPLIANCE QC22224

91 PASS BB = 96 PASS SURROGATES, %REC : TFT =

MATRIX SPIKE DUP: ACTUAL CALC % REC STATUS FILE 142J005 (

LUFT (ug/L) 2000 1666.96 83.3% IN COMPLIANCE QC22225

SURROGATES, %REC : TFT = 77 PASS BB = 82 PASS

AVERAGE SPIKE RECOVERY:

LUFT 86.6% RPD: 7.5 IN COMPLIANCE

RECOVERY CCV/LCS LUFT +/-10% DIFF MS/MSD LUFT +/-25% DIFF

LIMITS

SURROGATES TRIFLUOROTOLUENE (TFT): 69-120 %

BROMOBENZENE (BB): 70-122 %

CURTIS & TOMPKINS, LTD



TVH GASOLINE BY P&T GC/FID CALIBRATION FOR WATER/SOIL

CO4 LUFT

CALIBRATION DATE: JANUARY 25, 1996

SEQUENCE: JAN25

AMOUNT (ng)	SAMPLE AREA	FILE	Y CALC	RESIDUAL
500	618586.5	025J004	672.3	172.3
2500	2511783.8	025J00 5	2296.4	-203.6
5000	5551755	025J006	4904.1	- 95.9
10000	11680494	025J007	10161.5	161.5
15000	17316556	025J008	14996.2	-3.8
25000	28942862	025J009	24969.5	-30.5

Regression Output:

Constant	141.7110809
Std Err of Y Est	163.8457844
R Squared	0.999744835
No. of Observations	6
Degrees of Freedom	4

RANGE = 0



INSTRUMENT: GC04

SEQUENCE: MAY18

I.C.DATE: 01/25/96

DATE OF INJECTION18-MAY-

MATRIX: WATER

BATCH: 27666

QC22012

RESULT FILE: G:\GC04\139J001 LABORATORY CONTROL SAMPLE:

AMOUNT (ug/L) ACTUAL CALC % REC STATUS
LUFT 2000 2004.89 100.2% IN COMPLIANCE LUFT

SURROGATES, %REC: TFT = 100 PASS BB = 100 PASS

_____ RESULT FILE: G:\GC04\139J014 CONTINUING CALIBRATION CHECK:

AMOUNT (ug/L) ACTUAL CALC % REC STATUS

2000 1886.72 94.3% IN COMPLIANCE LUFT

SURROGATES, %REC: TFT = 105 PASS BB = 105 PASS

RESULT FILE: G:\GC04\139J029 CONTINUING CALIBRATION CHECK:

AMOUNT (ug/L) ACTUAL CALC % REC STATUS

2000 1986.48 99.3% IN COMPLIANCE LUFT

SURROGATES, %REC: TFT = 106 PASS BB = 106 PASS

RESULT FILE: G:\GC04\139J037 CONTINUING CALIBRATION CHECK:

AMOUNT (ug/L) ACTUAL CALC % REC STATUS

2000 1976.44 98.8% IN COMPLIANCE LUFT

SURROGATES, %REC: TFT = 106 PASS BB = 108 PASS

MSS sample: 125496-009 FILE 139J006

LUFT (ug/L)

SURROGATES, %REC: TFT = 92 PASS BB = 92 PASS _____

MATRIX SPIKE: ACTUAL CALC % REC STATUS 139J007 LUFT (ug/L) 2000 2074.57 103.7% IN COMPLIANCE QC22014

SURROGATES, %REC: TFT = 104 PASS BB = 103 PASS

MATRIX SPIKE DUP: ACTUAL

RIX SPIKE DUP: ACTUAL CALC % REC STATUS FILE 139J008 LUFT (ug/L) 2000 2069.11 103.5% IN COMPLIANCE QC22015

SURROGATES, %REC: TFT = 105 PASS BB = 105 PASS

AVERAGE SPIKE RECOVERY:

103.6% RPD: 0.3 IN COMPLIANCE LUFT

RECOVERY CCV/LCS LUFT +/-10% DIFF MS/MSD LUFT +/-25% DIFF

LIMITS

SURROGATES TRIFLUOROTOLUENE (TFT): 69-120 %

TVH2 - GC-04 RTX-1 FileName : G:\GC04\139J028.raw Date: 5/19/96 2:39 AM Page 1 of 1 Start Time : 0.00 min End Time : 17.00 min Low Point : 37.62 mV High Point: 287.62 mV Scale Factor: -1 Plot Offset: 38 mV Plot Scale: 250 mV Response [mV] 125615-001 0.88 1.17 -1.27 1.62 -2.00 2 36 TRIFLUOROT --2.88 3.09 3.34 3.60 7^{3.8}3 TOLUENE ETHYLBENZE = BROMOBENZE --6.22 -6.47 GAS PEAK 1 = $\frac{6.74}{6.90}$ GAS PEAK 3 -693⁴³ .90 8.09 9.26 .47 -9.94 -13.35

16.49

015

ਸ o

tron Time [min]

FileName : G:\GC04\142J006.raw

Start Time : 0.00 min

Scale Factor: -1

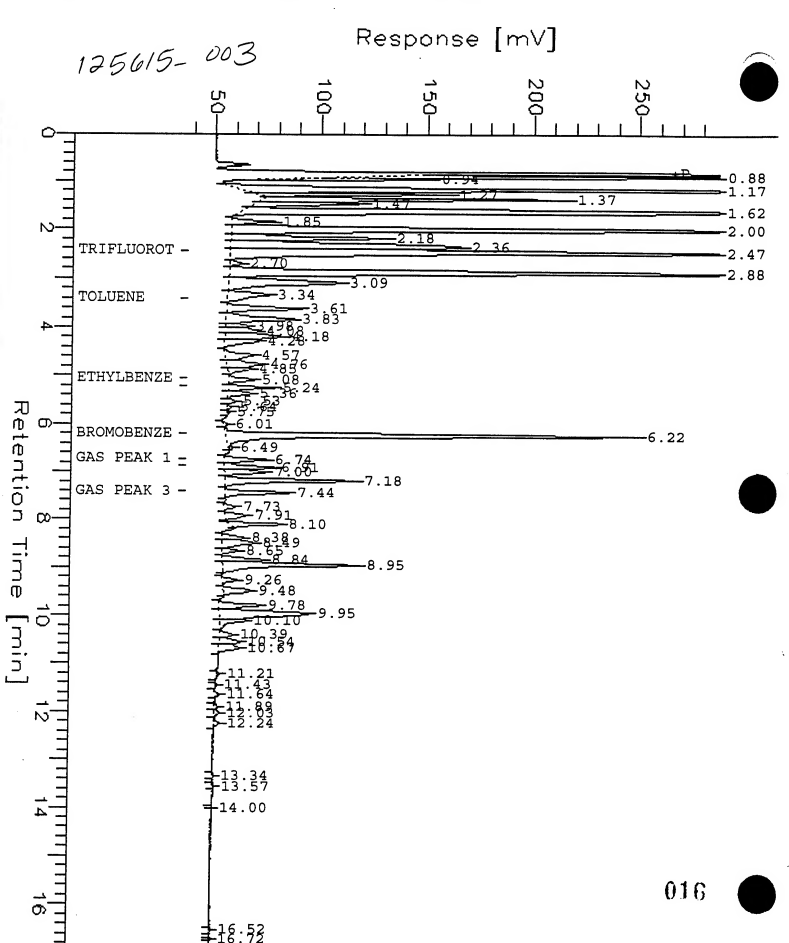
End Time : 17.00 min

Plot Offset: 37 mV

Date : 5/21/96 2:06 PM Low Point : 37.40 mV

Plot Scale: 250 mV

Page 1 of 1 High Point : 287.40 mV



Lab #: 125615

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

roject#: 722450.14020 Location: Beale AFB

Prep Method: EPA 5030

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ 05/08/96 Sample Date: Lab ID: 125496-009 05/08/96 Received Date: Matrix: Water Prep Date: 05/18/96 Batch#: 27666 Analysis Date: 05/18/96

Units: ug/L Diln Fac: 1

MS Lab ID: QC22014

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Gasoline	2000	<50.00	2075	104	75-125
Surrogate	%Rec	Limits			
Trifluorotoluene Bromobenzene	104 103	69-120 70-122			

MSD Lab ID: QC22015

alyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Gasoline	2000	2069	104	75-125	0	<20
Surrogate	%Rec	Limit	ts			
Trifluorotoluene Bromobenzene	105 105	69-12 70-12				

[#] Column to be used to flag recovery and RPD values with an asterisk

RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits

Lab #: 125615

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M) Prep Method: EPA 5030

Project#: 722450.14020

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Sample Date: 05/16/96 Lab ID: 125616-004 Received Date: 05/17/96 Matrix: Water Prep Date: 05/21/96 Analysis Date: 05/21/96

Batch#: 27719 Units: ug/L Diln Fac: 1

MS Lab ID: QC22224

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Gasoline	2000	<50.00	1796	90	75-125
Surrogate	%Rec	Limits			
Trifluorotoluene Bromobenzene	91 96	69-120 70-122			

MSD Lab ID: QC22225

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Gasoline	2000	1667	83	75-125	8	<20
Surrogate	%Rec	Limit	s			
Trifluorotoluene Bromobenzene	77 . 82	69-12 70-12	-			

Column to be used to flag recovery and RPD values with an asterisk

* Values outside of QC limits RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

Lab #: 125615

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc. Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Prep Method: EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

 Matrix:
 Water
 Prep Date:
 05/18/96

 Batch#:
 27666
 Analysis Date:
 05/18/96

Units: ug/L Diln Fac: 1

LCS Lab ID: QC22012

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	2005	2000	100	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	100	69-120		
Bromobenzene	100	70-122		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

^{&#}x27;pike Recovery: 0 out of 1 outside limits



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Water

Prep Method: EPA 5030

Location: Beale AFB

LABORATORY CONTROL SAMPLE

Prep Date: 05/21/96

Analysis Date: 05/21/96

Batch#: 27719 Units: ug/L Diln Fac: 1

Matrix:

LCS Lab ID: QC22221

Analyte	Result	Spike Added	%Rec #	Limits	
Gasoline	1802	2006	90	80-120	•
Surrogate	%Rec	Limits			
Trifluorotoluene Bromobenzene	92 94	69-120 70-122			

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

Spike Recovery: 0 out of 1 outside limits

Lab #: 125615

Page 1 of 1

TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc. roject#: 722450.14020

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 5030

Location: Beale AFB

METHOD BLANK

Matrix: Water Batch#: 27666 Units: ug/L

Diln Fac: 1

05/18/96 Prep Date: Analysis Date: 05/18/96

MB Lab ID: QC22011

Analyte	Result	3
Gasoline	<50	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene	99	69-120
Bromobenzene	104	70-122

Lab #: 125615 BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

METHOD BLANK

Matrix: Water Batch#: 27719 Units: ug/L Diln Fac: 1

Prep Date: 05/21/96 Analysis Date: 05/21/96

MB Lab ID: QC22223

Analyte	Result	
Gasoline	<50	
Surrogate	%Rec ·	Recovery Limits
Trifluorotoluene	84	69-120
Bromobenzene	82	70-122

TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)
Prep Method: EPA 5030

Project#: 722450.14020 Location: Beale AFB

Sample #	Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
125615-001	BE-MP10-01	27666	05/14/96	05/19/96	05/19/96	
125615-002	BE-MP12-01	27666	05/15/96	05/19/96	05/19/96	
125615-003	BE-MP13-01	27719	05/14/96	05/21/96	05/21/96	
125615-004	BE-MP47-01	27666	05/14/96	05/19/96	05/19/96	

Matrix: Water

Analyte Diln Fac:	Units	125615-001 1	125615-002 1	125615-003 5	125615-004 1
Gasoline	ug/L	1400	<50	5900	<50
Surrogate					
ifluorotoluene	%REC	96	89	90	92
Bromobenzene	%REC	117	89	98	93



TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
125615-005 TRIP BLANK	27666	05/15/96	05/19/96	05/19/96	

Matrix: Water

Analyte Diln Fac:	Units	125615-005 1		
Gasoline	ug/L	<50		
Surrogate			 	
Trifluorotoluene	%REC	92		
Bromobenzene	%REC	93		

Benefis



Curtis & Tompkins, Ltd

COOLER RECEIPT CHECKLIST

Logi:		4(-B	
Ciicii	1.0,000		***
A.	Preliminary Examination Phase		
	Date Opened: Ship By (print): J. William (sign)	سس	$\widehat{}$
1.	Date Opened: St. By (print): J. W. (sign) Did cooler come with a shipping slip (airbill, etc.)? LEVES enter comics name and cirbill number: Teal (sign)	V ES	NO
	If YES, enter carrier name and anom number.	U)	
2.	Were custody seals on outside of cooler?	YES	100
	How many and where? Seal date: Seal name:	N	12
3.	Were custody seals on outside of cooler? How many and where? Seal date: Seal name: Were custody seals unbroken and intact at the date and time of arrival?	. YES	NON'
4.	Were custody papers dry and intact when received?	(YES	NO
5.	Were custody papers filled out properly (ink, signed, etc.)?	VES	NO
6.	Did you sign the custody papers in the appropriate place?	(E S	NO
7 .	Was project identifiable from custody papers?	KES	NO
	If YES, enter project name at the top of this form.	1 –	
8.	If required, was sufficient ice used?	CES	NO
	Type of ice: Temperature: 40 · C		
	,	` .	
B.	Login Phase		
	Date Logged In: 5/7 By (print): dull (sign)	<u>Sure</u>	<u>دب</u>
1.	Describe type of packing in cooler: bull usep Did all bottles arrive unbroken?	····	
2.		. ŒS	
3.	Were labels in good condition and complete (ID, date, time, signature, etc.)?.		
4.	Did bottle labels agree with custody papers?	. OFER	NO
5.	Were appropriate containers used for the tests indicated?	XES	NO
6.	Were correct preservatives added to samples?		
7.	Was sufficient amount of sample sent for tests indicated?	1595	NO
8.	Were bubbles absent in VOA samples? If NO, list sample Ids below	E S	NO
9.	Was the client contacted concerning this sample delivery?	YES	NO
	If YES, give details below.		
	Who was called? By whom? Date	e:	
Addit	ional Comments:		
			
			
			
			
Filenam	e: F:\qc\forms\cooler.wpd R	ev. 1 4/95	



TPH/Purgeable Data



Chain of Custody

USA CHAIN OF CUSTODY RECORD

10F

PAGE

SOI Marina Village PKWY, Suite 200 Alameda, California 94501

FARSONS ENGINEERING SCIENCE, INC.

Phone: (510) 769-0100 FAX: (510) 769-9244

handbook requirements. **according to AFCEE** and holding times Note: All analyses Remarks **OTHER** INORGANICS ORGANICS \$6107 863 (0508/0502) \$EMT by X318 e101 663 (172 108/0502) × MALYTES (METHOD) ソ 7 NO. OF CONTAINERS water water Water water water water water Matrix Depth ١ Begin Depth Project Manager: Todd Wiedemeier (Derwer, CO) Project Name/Location: Beale AFB, UST Site 15/14/96 11245 Site Manager: Michael Phelps (Alameda, CA) SUM 196 1305 550/ SHEPPE 1100 9501 96/51/5 Sampler(s): (Initials and Signature(s))
Michael B. Phelos 2/14/146 2/12/18 Date Project No.: 722450.14020 E. -4 BE-MP47-01 BE-MP19-01 7 BE- MP15-DI BE-MPIZ-BI Togo Blank Sample ID

Distribution: Original accompanies shipment: photocopy kent by samplers: copy FAXed to Persons ES Site Manager by Jahoratory upon receint of camples holding time. Expedite tury

MANUEL C

4. TMBs: 1,2,3-trimethylbenzene, 1,2,4-trimethylbenzene, and 1,3,5-trimethylbenzene.

Secondary column confirmation required.
 Quantify all carbon ranges: diesel, jet fuel, cil/grease.
 Delonized water extraction.

REMARKS/COMMENTS:

RECEIVED FOR LAB BY: (SIGNATURE) DATE

TIME

DATE

RELINGUISHED BY: (SIGNATURE)

RECEIVED BY: (SIGNATURE)

630

5/1c/96

DATE

(SIGNATURE)

water

water

water

water water water Resampled due to laboratory-missed



Curtis & Tompkins, Ltd., Analytical Laboratories, Since 1878

2323 Fifth Street, Berkeley, CA 9471O, Phone (510) 486-0900

COVER PAGE

Laboratory Number 125615

Parsons Engineering Science, Inc.

Project#: 722450.14020

1301 Marina Village Parkway

Location: Beale AFB

Suite 200

Alameda, CA 94501

Sample ID	Lab ID
BE-MP10-01	125615-001
BE-MP12-01	125615-002
BE-MP13-01	125615-003
BE-MP47-01	125615-004
TRTP BLANK	125615-005

NOTE -

I certify that this data package has been reviewed for technical correctness and completeness. Please see attached narrative for a discussion of any analytical problems related to this sample set. Release of this data has been authorized by the Laboratory Manager or the Manager's designee, as verified by the following signatures.

Signature:

Title: Operations

Date:

Signature:

Tuesa KMOrrason

Date:

= 5/23/94

Title: Project Manager



Laboratory Number: 125615

Client: Parsons Engineering Science

Project#: 722450.14020 Location: Beale AFB Sample Date: 05/14,15/96 Receipt Date: 05/17/96



CASE NARRATIVE

This hardcopy data package contains sample results and batch QC for four water samples and one trip blank received from the above referenced project on May 17, 1996. All samples were received cold and intact.

TPH/Purgeable: No analytical problems were encountered.

BTXE (EPA 8020): Because TPH/gasoline and BTXE are analyzed simultaneously, only one type of matrix spikes are performed. The matrix spikes associated with these samples were gasoline. Both laboratory control samples passed acceptance criteria and no analytical problems were encountered.

Calla

COOLER RECEIPT CHECKLIST

Curtis	&	Tompkins.	Lid.

Login#	Date Received: 5/15 Number of Coolers:	1	
Client:			
Chem.	110,000.		
A.	Preliminary Examination Phase Date Opened: Strong By (print): Strong (sign)	عدر	2
1.	Did cooler come with a shipping slip (airbill, etc.)?	. YES	<u>র্</u>
2.	Were custody seals on outside of cooler?	. YES	NO
	How many and where? Seal date: Seal name:		At
3.	Were custody seals unbroken and intact at the date and time of arrival?	YES	NO .
4.	Were custody papers dry and intact when received?		
5.	Were custody papers filled out properly (ink, signed, etc.)?		
6.	Did you sign the custody papers in the appropriate place?	_	
7.	Was project identifiable from custody papers?		
••	If YES, enter project name at the top of this form.	. ==	
8.	If required was sufficient ice used?	YES	NO
o .	If required, was sufficient ice used? Type of ice: Temperature: H.45"	. عود	1.0
	Typo or ice.		
B.	Login Phase Date Logged In: 3/16 By (print): 0.00 (sign) (sign)	الايد	
1.	Describe type of packing in cooler:		
2.	Did all bottles arrive unbroken?	(YES	NO
3.	Were labels in good condition and complete (ID, date, time, signature, etc.)?	. XES	NO
4.	Did bottle labels agree with custody papers?		
5.	Were appropriate containers used for the tests indicated?		
6.	Were correct preservatives added to samples?	_	
7.	Was sufficient amount of sample sent for tests indicated?	/1	
8.	Were bubbles absent in VOA samples? If NO, list sample Ids below	1 , 1	
9.	Was the client contacted concerning this sample delivery?	_	
	If YES, give details below.		\mathcal{O}
	Who was called? By whom? Date:		
Additic	onal Comments:		
Filename:	F:\qc\forms\cooler.wpd Rev	. 1 4/95	



BATCH QC REPORT

Page 1 of 1

BTXE

lient: Parsons Engineering Science, Inc.

Analysis Method: EPA 8020

EPA 5030

Project#: 722450.14020 Location: Beale AFB

Prep Method:

LABORATORY CONTROL SAMPLE

Water

Prep Date:

03/29/96

Batch#: 26707 Units: ug/L

Diln Fac: 1

Matrix:

Analysis Date: 03/29/96

LCS Lab ID: QC18179

Analyte	Result	Spike Added	%Rec #	Limits
Benzene	20.8	20	104	80-120
Toluene	21.3	20	107	80-120
Ethylbenzene	19.9	20	100	80-120
m,p-Xylenes	42.9	40	107	80-120
o-Xylene	20.9	20	105	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene	97	58-130		
Bromobenzene	86	62-131		

Column to be used to flag recovery and RPD values with an asterisk lates outside of QC limits ke Recovery: 0 out of 5 outside limits

Fax: (510) 769-9244 Engineering-Science, Inc. 1301 Marina Village Parkway, Suite 200 Alameda, California 94501 Phone: (510) 769-0100 Fax: (510) 76

CHAIN OF CUSTODY RECORD 678 421

P.

PAGE

RECEIVED BY: (SIGNATURE) REMARKS SIMIT ONNO TIME TO BE COMPOSITED BY LAB TIME ANALYSIS REQUIRED DATE REMARKS: RELINQUISHED BY(SIGNATURE) TIME Q-HOL DRESERVED DATE X X X X RECEIVED FOR LABORATORY BY: (SIGNATURE) メ X RECEIVED BY: (SIGNATURE) 722420.14022 NO. OF CONTAINERS + 7 7 7 SAMPLE LOCATION ER PROJ. #: D. FCK t 3 • Drainege PROJECT MANAGER: TIME 13/15/26 1830 TIME M. Phelps 7 DATE DATE + Parky 本の Beak AFB UST RELINQUISHED BY: (SIGNATURE) **BLINQUISHED BY: (SIGNATURE)** Marian J. Prein PROJECT NAME/LOCATION: TIME 1230 1415 SAMPLER(S): (SIGNATURE) 0271 BE-SWOI-01 3/15/96 1120 Marcus Pierce DATE C & 1 LABORATORY: BE-SWIZ-PI BE-SW03-DI BE-SW02-D SAMPLE ٦

MISTRIBUTION: WIITE: ACCOMPANIES SHIPMENT & RET'N WITH LAB REPORT, CANARY: LAB COPY, PINK. FIELD COPY

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Page 1 of 1

BTXE

lient: Parsons Engineering Science, Inc.

Project#: 722450.14020 Location: Beale AFB Analysis Method: EPA 8020

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124848-001 BE-SW01-01	26707	03/15/96	03/29/96	03/29/96	
124848-002 BE-SW02-01	26707	03/15/96	03/29/96	03/29/96	
124848-003 BE-SW12-01	26707	03/15/96	03/29/96	03/29/96	
124848-004 BE-SW03-01	26707	03/15/96	03/29/96	03/29/96	

Analyte [Diln Fac:	Inits	124848-001 1	124848-002 1	124848-003 1	124848-004 1
Benzene	ug/L	<0.5	<0.5	<0.5	<0.5
Toluene	ug/L	<0.5	<0.5	<0.5	<0.5
Ethylbenzene	ug/L	<0.5	<0.5	<0.5	<0.5
m,p-Xylenes	ug/L	<0.5	<0.5	<0.5	<0.5
o-Xylene	ug/L	<0.5	<0.5	<0.5	<0.5
,2,3-Trimethylbenzene		<0.5	<0.5	<0.5	<0.5
,2,4-Trimethylbenzene	<i>-</i> ,	<0.5	<0.5	<0.5	<0.5
1,3,5-Trimethylbenzene	ug/L	<0.5	<0.5	<0.5	<0.5
Surrogate					
Trifluorotoluene	%REC	94	95	94	95
Bromobenzene	%REC	82	82	80	82



BATCH QC REPORT

Page 1 of 1

BTXE

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: EPA 8020

Prep Method: EPA 5030

METHOD BLANK

 Matrix:
 Water
 Prep Date:
 03/29/96

 Batch#:
 26707
 Analysis Date:
 03/29/96

Units: ug/L Diln Fac: 1

MB Lab ID: QC18177

Analyte	Result	
Benzene	<0.5	
Toluene	<0.5	
Ethylbenzene	<0.5	
m,p-Xylenes	<0.5	
o-Xylene	<0.5	
1,2,3-Trimethylbenzene	<0.5	
1,2,4-Trimethylbenzene	<0.5	
1,3,5-Trimethylbenzene	<0.5	
Surrogate	%Rec	Recovery Limits

Surrogate	%Rec	Recovery Limits
Trifluorotoluene	95	58-130
Bromobenzene	83	62-131



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

client: Parsons Engineering Science, Inc. Analysis Method:

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M)

Prep Method: EPA 5030

LABORATORY CONTROL SAMPLE

Matrix: Water Batch#: 26707 Units: ug/L

Prep Date: 03/28/96 Analysis Date: 03/28/96

Diln Fac: 1

LCS Lab ID: QC18178

Analyte	Result	Spike Added	%Rec #	Limits
Gasoline	1938	2000	97	80-120
Surrogate	%Rec	Limits		
Trifluorotoluene Bromobenzene	102 106	69-120 70-122		

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

^{&#}x27;pike Recovery: 0 out of 1 outside limits



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M

Project#: 722450.14020

Prep Method: EPA 5030

Location: Beale AFB

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

Field ID: ZZZZZZ Lab ID: 124892

ZZZZZZ Sample Date: 03/15/96 124892-016 Received Date: 03/19/96 Water Prep Date: 03/28/96

Matrix: Water Batch#: 26707 Units: ug/L Diln Fac: 1

Prep Date: 03/28/96 Analysis Date: 03/28/96

MS Lab ID: QC18304

Analyte	Spike Added	Sample	MS	%Rec #	Limits
Gasoline	2000	<50.00	1885	94	75-125
Surrogate	%Rec	Limits			
Trifluorotoluene Bromobenzene	102 105	69-120 70-122			

MSD Lab ID: QC18305

Analyte	Spike Added	MSD	%Rec #	Limits	RPD #	Limit
Gasoline	2000	1841	92	75-125	2	<20
Surrogate	%Rec	Limit	:8			
Trifluorotoluene Bromobenzene	102 105	69-12 70-12	_	• , , , , ,		

[#] Column to be used to flag recovery and RPD values with an asterisk

RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

^{*} Values outside of QC limits



TVH-Total Volatile Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020 Location: Beale AFB

Prep Method: EPA 5030

Sample # Client ID	Batch #	Sampled	Extracted	Analyzed	Moisture
124848-001 BE-SW01-01	26707	03/15/96	03/29/96	03/29/96	
124848-002 BE-SW02-01	26707	03/15/96	03/29/96	03/29/96	
124848-003 BE-SW12-01	26707	03/15/96	03/29/96	03/29/96	
124848-004 BE-SW03-01	26707	03/15/96	03/29/96	03/29/96	

Analyte Diln Fac:	Units	124848-001 1	124848-002 1	124848-003 1	124848-004 1
Gasoline	ug/L	<50	<50	<50	<50
Surrogate					
Trifluorotoluene	%REC	97	97	96	98
comobenzene	%REC	89	89	87	89



BATCH QC REPORT

Page 1 of 1

TVH-Total Volatile Hydrocarbons

Client: Parsons Engineering Science, Inc.

Project#: 722450.14020

Location: Beale AFB

Analysis Method: CA LUFT (EPA 8015M Prep Method:

EPA 5030

METHOD BLANK

Matrix: Water Batch#: 26707 Units: ug/L

Diln Fac: 1

Prep Date: Analysis Date:

03/29/96 03/29/96

MB Lab ID: QC18177

Analyte	Result	
Gasoline	<50	
Surrogate	%Rec	Recovery Limits
Trifluorotoluene Bromobenzene	96 89	69-120 70-122

iesel

GC15 Channel A TEH

Sample Name : CCV,96WS1948,DSL FileName : C:\GC15\CHB\080B047.RAW

Method

: BTEH.MTH Start Time : 0.01 min

Nie Factor: 0.0 End Time : 31.91 min

Plot Offset: 44 mV

Sample #: 500MG/L

Date: 3/25/96 05:16 PM

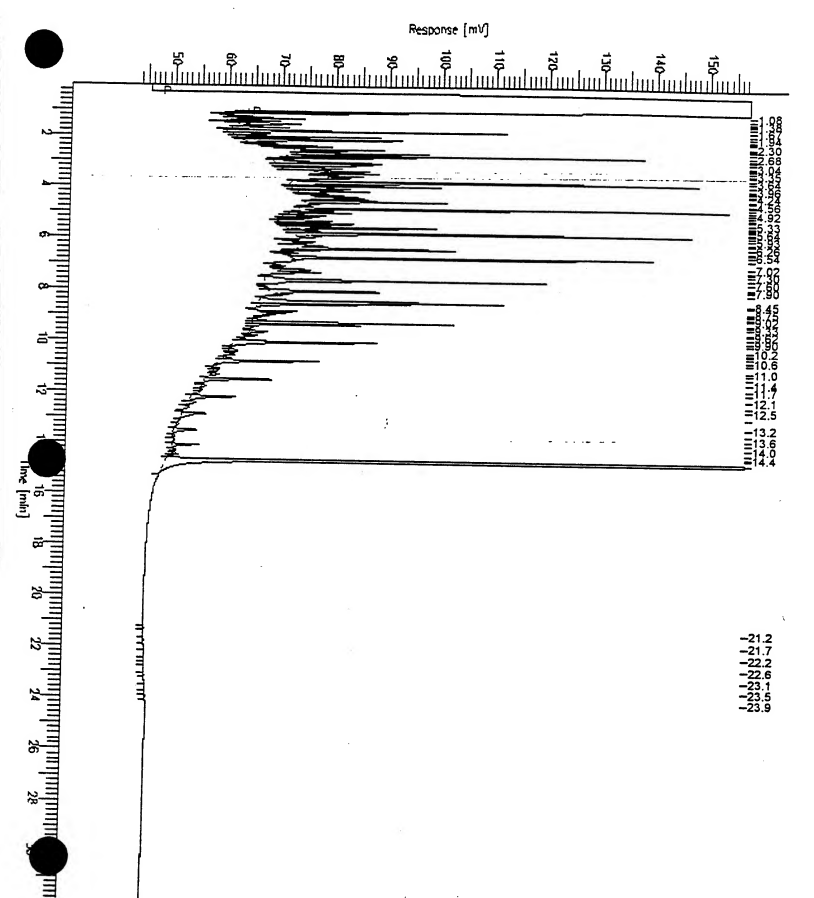
Time of Injection: 3/22/96 01:48 AM

Low Point: 43.61 mV

High Point : 157.43 mV

Page 1 of 1

Plot Scale: 113.8 mV



Motor OU

Sample Name : CCV, 95WS1596, MO

FileName : C:\GC15\CHB\080B051.RAW

: BTEH.MTH Methoa

Start Time : 0.01 min 0.0 Scale Factor:

End Time : 31.91 min Plot Offset: 41 mV

Sample #: 500MG/L

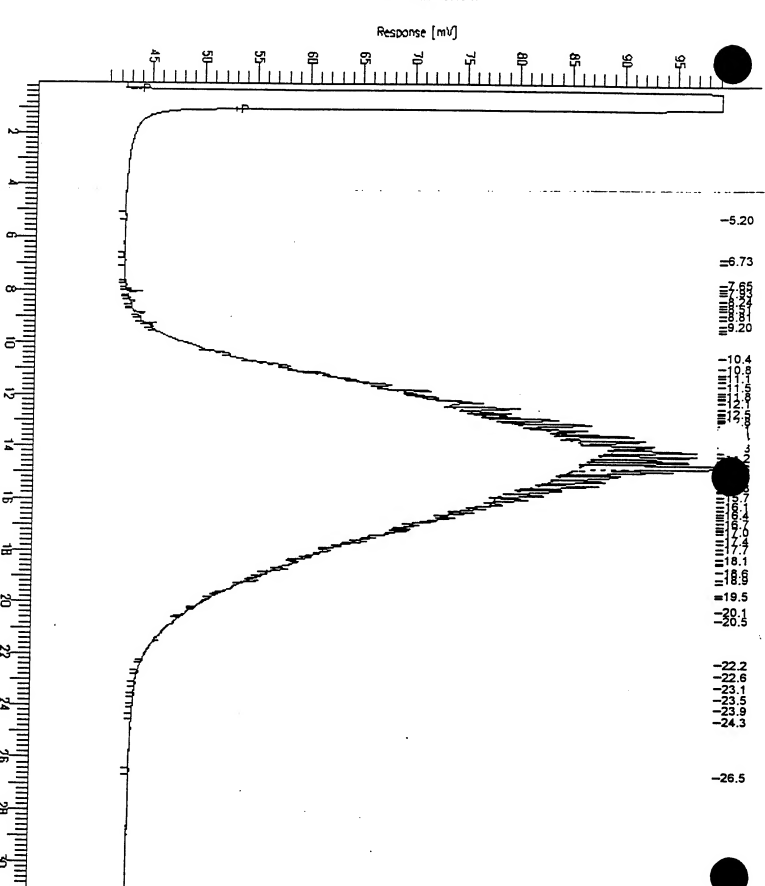
Date: 3/25/96 05:14 PM Time of Injection: 3/22/96 04:44 AM

Low Point : 40.69 mV

High Point : 99.19 mV

Page 1 of 1

Plot Scale: 58.5 mV



Dullogace

Sample Name: S,124848-003,26513 FileName : C:\GC15\CHB\080B055.raw

Method : DUAL

Start Time : 0.00 min Scale Factor: 0.0

End Time : 31.90 min Plot Offset: 32 mV

Sample #: 500:2.5

Date: 3/22/96 08:13 AM

Time of Injection: 3/22/96 07:39 AM

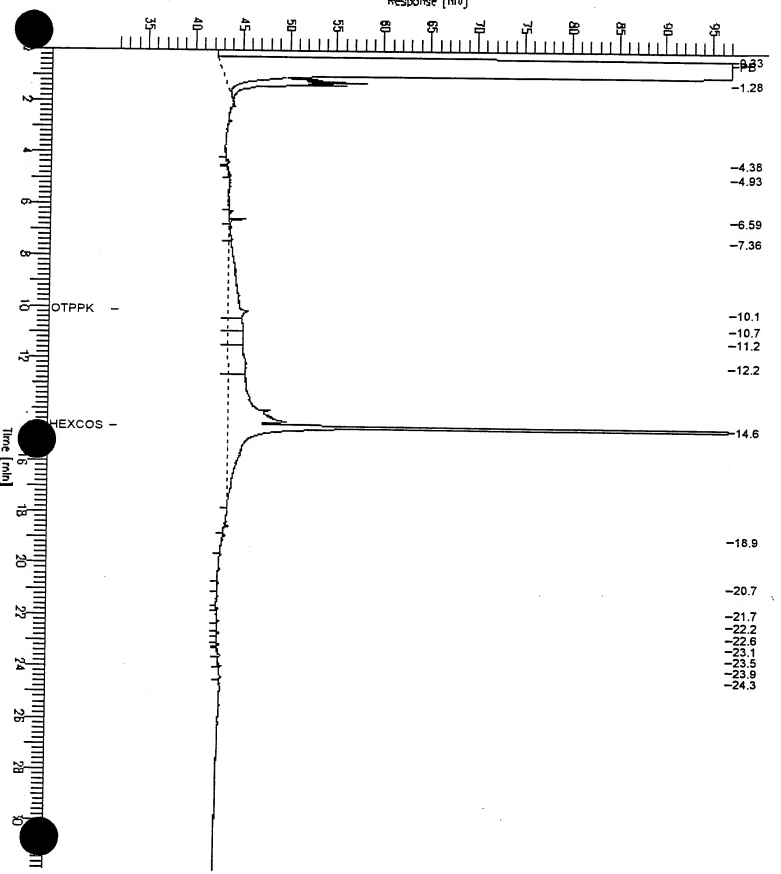
Low Point : 32.00 mV

High Point : 97.00 mV

Page 1 of 1

Plot Scale: 65.0 mV





Channel A TEH GCID

Sample Name : CCV,96WS2068,JP5 FileName : C:\GC15\CHB\080B050.RAW

Method : BTEH.MTH

Start Time : 0.01 min Scale Factor: 0.0

End Time : 31.91 min

Plot Offset: 43 mV

Sample #: 250MG/L

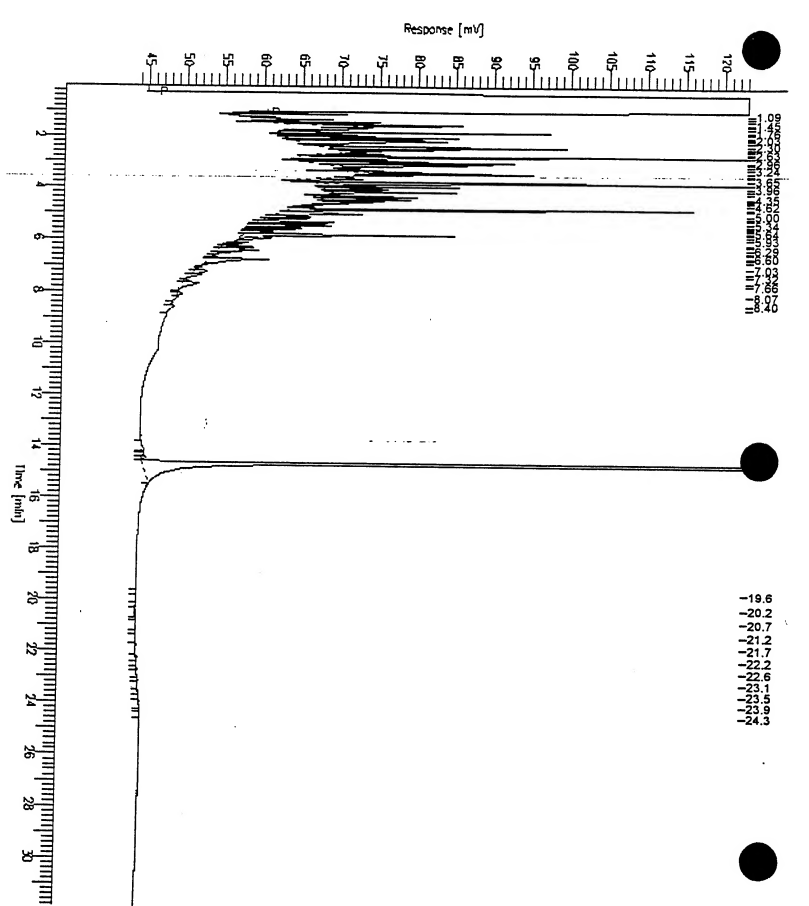
Date: 3/25/96 05:13 PM

Time of Injection: 3/22/96 04:00 AM Low Point : 43.16 mV

High Point : 123.07 mV

Page 1 of 1

Plot Scale: 79.9 mV



BATCH QC REPORT

Page 1 of 1

TEH-Tot Ext Hydrocarbons

lient: Parsons Engineering Science, Inc.

Analysis Method: CA LUFT (EPA 8015M)

Project#: 722450.14020

Prep Method: EPA 3520

Location: Beale AFB

BLANK SPIKE/BLANK SPIKE DUPLICATE

Prep Date:

03/19/96

Matrix: Water Batch#: 26513 Units: ug/L

Analysis Date: 03/21/96

Diln Fac: 1

BS Lab ID: QC17467

Analyte	Spike Added BS	%Rec #	Limits
Diesel Range	2475 2218	90	60-140
Surrogate	%Rec Limits	S	
Hexacosane	96 60-140	0	The state of the s

BSD Lab ID: QC17468

Analyte	Spike Added	BSD	%Rec #	Limits	RPD #	Limit
iesel Range	2475	2330	94	60-140	5	<35
surrogate	%Rec	Limits				
Hexacosane	95	60-14	0			

[#] Column to be used to flag recovery and RPD values with an asterisk

^{*} Values outside of QC limits

RPD: 0 out of 1 outside limits

Spike Recovery: 0 out of 2 outside limits

Sample Name: S,124848-002,26513 FileName : C:\GC15\CHB\080B054.raw

Method : DUAL

Start Time : 0.00 min Scale Factor: 0.0

End Time : 31.90 min Plot Offset: 32 mV

Sample #: 500:2.5

Date: 3/22/96 07:29 AM

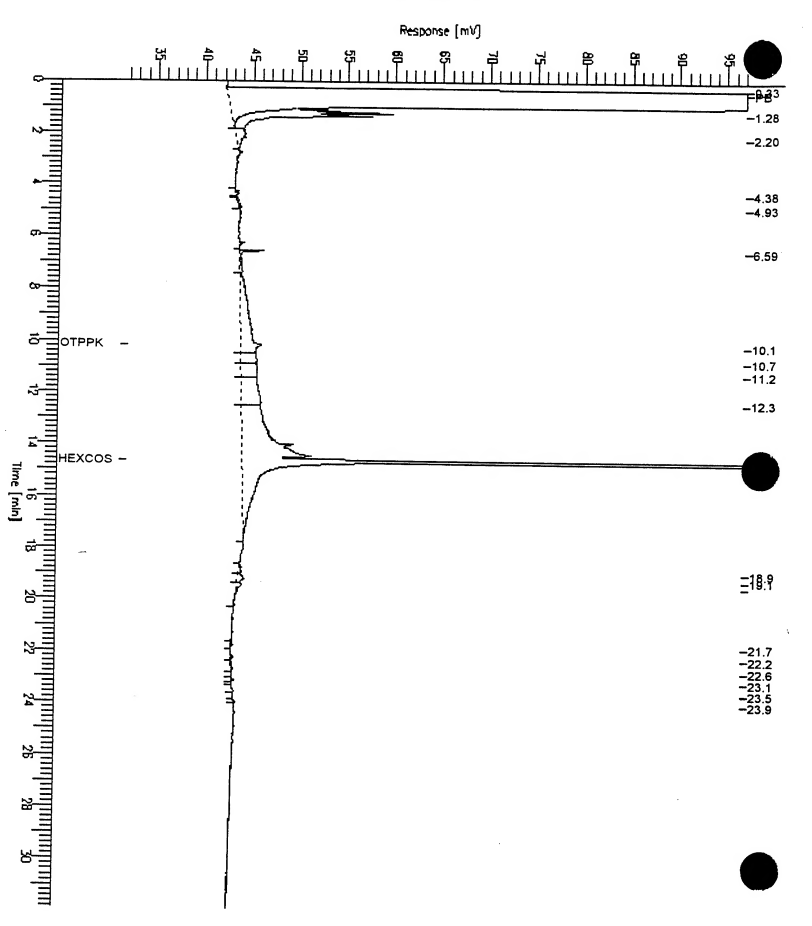
Time of Injection: 3/22/96 06:55 AM

Low Point : 32.00 mV

High Point : 97.00 mV

Page 1 of 1

Plot Scale: 65.0 mV



APPENDIX D

MODEL INPUT PARAMETERS AND HYDRAULIC TESTING RESULTS

CALIBRATED MODEL BOUNDARY CONDITIONS AND TRANSMISSIVITIES

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

1	2		3 	4 	5	6 	7 	8 	9	10	11	12	13	14 1	5 	16	17 18	19
Ш		Ш			ЩШ			ШШ								!!! !!!		
	129.75	129.6	7 129.5	8 129.	50 A	Α	Α	Α	Α	Α	Α	Α	В	В	С	С	С	126.09
	129.71	А	Α	Α	Α	Α	Ä	Α	Α	Α	Α	A	В	В	С	С	_с	125.98
	1	А	Α	Α	Α	Α	Α	Α	Α	Α	Α	В	В	В	В	В	С	125.86
	129 64	Α	Α	Α	Α	Α	Α	Α	Α	Α	Α	В	В	В	В	В	С	125.75
		А	Α	Α	Α.	Α	Α	Α	Α	Α	В	В	В	В	В	В	c	125.64
	129.57	Α	Α	Α	Α	Α	Α	Α	Α	В	В	В	В	В	С	С	С	125.59
	123.34	A	Α	Α	Α	Α	Α	Α	_A_	В	В	В	В	С	С	С	С	125.54
	129.50	A	Α	Α	Α	Α	Α	Α	В	В	В	В	В	С	С	С	С	125.48
	Α	Α	Α	Α	Α	Α	Α	В	В	В	В	В	В	С	С	С	С	125.43
	A	Α	Α	Α	Α	Α	В	В	В	В	В	В	В	С	С	С	С	125.38
	А	Α	Α	Α	Α	В	В	В	В	В	В	В	В	С	С	С	c	125.32
	А	Α	Α	Α	В	В	В	В	В	В	В	В	В	В	В	В	С	125.27
	А	Α	Α	В	В	В	В	В	В	В	В	В	В	В	В	В	С	125.21
	A	Α	В	В	В	В	В	В	В	В	В	В	В	В	В	В	С	125.16
	В	В	В	В	В	В	В	В	В	В	В	В	В	В	В	В	С	125.11
	В	В	В	В	В	В	В	В	В	В	В	В	В	В	В	В	С	125.05
	В	В	В	В	В	В	В	В	В	В	В	В	В	В	В	В	С	125.00
	С	С	С	С	В	В	В	В	В	В	В	В	В	В	В	В	С	124.95
	С	С	С	С	C	С	С	С	С	С	С	С	С	С	С	С	С	124.90
	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	124.85
++	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	124.80
	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	, C	124.75
	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	124.75
	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	124.70
	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	124.65
	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	124.60
	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	124.55
1	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	С	124.50
Ш			Ш															

not to scale; grid dimensions are square (20 feet by 20 feet)

A: Transmissivity = 8.1×10^{-4} ft/s B: Transmissivity = 3.96×10^{-4} ft/s C: Transmissivity = 4.5×10^{-5} ft/s

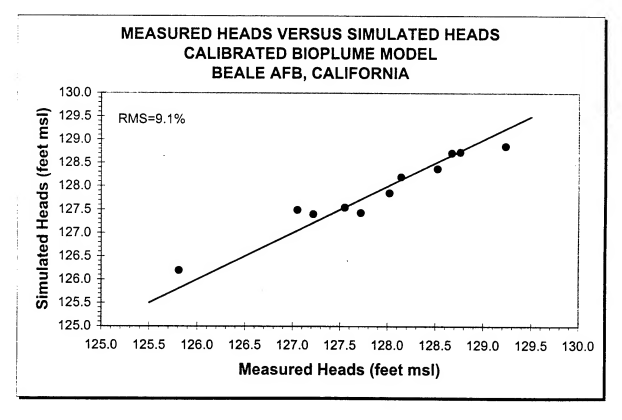


Constant Head cell (feet)
No flow cell

ROOT-MEAN-SQUARE ERROR FOR THE CALIBRATED FLOW MODEL

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

l 4:			h _m , Measured Head	h _s , Simulated Head	$(h_{\rm m} - h_{\rm s})^2$
Location	x-grid	y-grid	(feet msl)	(ft msl)	(ft ²)
MW-1	7	12	129.23	128.86	0.137
MP-9	9	11	128.67	128.71	0.002
MP-10	12	13	128.14	128.20	0.003
MP-11	17	5	128.02	127.86	0.027
MP-12	16	10	127.05	127.49	0.197
MP-13	16	14	127.22	127.40	0.034
MP-14	14	18	127.55	127.55	0.000
MP-15	11	22	127.72	127.43	0.082
MP-17	15	24	125.81	126.20	0.149
MP-21	3	17	128.76	128.73	0.001
MP-22	8	17	128.52	128.37	0.021
				RMS Error (ft) =	0.243
Differen	ice betweei	n minimun	n and maximum s	simulated heads (ft) =	2.66
				RMS Error (%)=	9.1%



EXAMPLE SOURCE TERM LCULATION FOR RESIDUAL LNAPL

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

Current Conditions (MP-10)

	Cs. Conc.		% Analyte in	ر ن	ڻ	K _w , Fuel-	ڻُ	Leaching
	Measured		Fresh	Conc. in Fresh	Leachable		Concentration	Rate To
	in Soil	% Analyte	Gasoline	Gasoline	Conc.	_	in Recharge	Groundwater
Contaminant	(mg/kg)	in TPH (%)	(%)	(mg/L)	(mg/L)	Coefficient	(mg/L) (n	(mg/day-ft²)
Benzene	0.0	0.0	1.8	15,928	0.0	291	0.0	0.0
Toluene	0.0	0.0	7.1	61,723	0.0	1,073	0.0	0.0
Ethylbenzene	0.0	0.0	1.8	15,648	0.0	3,956	0.0	0.0
Xylenes	2.2	0.282	9.0	77,858	2450	3,665	0.67	0.00479
TOTAL								0.00479

Calculated Initial Conditions (MP-10)

BTEX Concentration: BTEX Concentration:

Mass Loading:

1.92 mg/day-cell 0.16 mg/l-cell 157 ug/l-cell

	Cs. Conc. Measured in Soil	% Analyte	% Analyte in Fresh Gasoline	Conc. in Fresh Leachable	C _{fi} . Leachable	K _w , Fuel- Water Partition	Cw. Concentration in Recharge	Leaching Rate To Groundwater
Contaminant	_	in TPH (%)		(mg/L)	(mg/L)	Coefficient	(mg/L)	(mg/day-ft²)
Benzene	17.8	2.3	1.8	15,928	19,786	291	68.1	0.488
Toluene	0.69	8.8	7.1	61,723	76,673	1,073	71.5	0.513
Ethylbenzene	17.5	2.2	1.8	15,648	19,437	3,956	4.9	0.035
Xylenes	86.9	11.1	9.0	77,858	96,716	3,665	26.4	0.189
TOTAL								1.23

490 mg/day-cell 40.1 mg/l-cell 40,102 ug/l-cell BTEX Concentration: BTEX Concentration:

Mass Loading:

Calculated First-Order Decay Rate
0.000447 per day

0.16 per year

EXAMPLE SOURCE TERM LCULATION FOR RESIDUAL LNAPL

(continued)

UST Sites 10-494, 10-495, and 10-496 Intrinsic Remediation TS Beale AFB, California

NOTES AND ASSUMPTIONS

Properties of Gasoline	ine	Source:	Source: Wiedemeier et al. (1995)	et al. (1995)	
	density				
	(g/m³)	K _{fw} (-)	F _r (-)	C _r (mg/L)	C _w (mg/L)
Gasoline	720,000	-			
Benzene	868,000	291	1.8	15,928	54.8
Toluene	866,900	1,073	7.1	61,723	57.6
Ethylbenzene	866,900	3,956	1.8	15,648	4.0
Xylenes	868,467	3,665	0.6	77,858	21.2
TOTAL	٠		19.7		

[analytical result at MP-10] 780 mg/kg 969 mg/kg 1.11 in/yr 2.53E-04 ft/day current TPH in soil Initial TPH in soil

recharge rate recharge rate cell area constant injection rate

400 ft2/cell 28.3 L/ft3 5.00E-06 ft^3/s

=(780-2.2)/(1-0.197)

[5% of annual rainfall]

 K_{tw} = fuel-water partition coefficient F_{m} = mass fraction of compound in gasoline $C_f(g/m^3) = Concentration in fuel <math>C_{sat}(g/m^3) = Concentration in water$

				AQTESOLU
g Science			C: TIMATES: 51 ft ² /sec	٠
сонтыну: Parsons Engineering Science	.14	Test #1)	DATA SET: AM11R.DAT 10/31/96 AQUIFER MODEL: Confined SOLUTION METHOD: Cooper et al. TEST DATA: H0= 2.114 ft r_= 0.083 ft r_= 0.344 ft r_= 0.344 ft T = 0.0002351 ft ² /s S = 1.E-05	
: Parson	PROJECT: 722450,14			
COMPANY	PROJECT	Head		-
		(Rising	10.001	
	₩.			lime (sec)
	Beale AFB, CA	MW-1		
	1		0.01	
· AFCEE	LOCATION: UST Site		0.6 	
CL IENT:	LOCATI		Displacement, H/HO (ft)	

•

93			ວູ	
ngineering Scien		#5)	DATA SET: AM12R.DAI 10/31/96 AQUIFER MODEL: Confined SOLUTION METHOD: Cooper et al. TEST DATA: H0= 1.516 ft r_c= 0.083 ft r_w= 0.344 ft PARAMETER ESTIMATES: I = 0.0002924 ft²/sec S = 1.E-05	
сомрану: Parsons Engineering Science	PROJECT: 722450,14	Head Test #2)		
5		(Rising H	10. 100.	
	Beale AFB, CA	MW-1 (
CLIENT: AFCEE	LOCATION: UST Site		0.8	

CLIENT: AFCEE LOCATION: UST Site Beale AFB, CA	COMPANY: Parsons E. PROJECT: 722450,14	сонгену: Parsons Engineering Science гволест: 722450.14
MW-2 (Falling	Head Te	Test #1)
10. <u>ETTTTTTTTTTT</u>	<u> </u>	DATA SET: AM21F.DAT 10/31/96
-	11111	AQUIFER MODEL: Confined SOLUTION METHOD: Bouwer-Rice
Gocement (ft)	huurra — huu	TEST DATA: H0= 3.039 ft r _c = 0.083 ft r _w = 0.344 ft L = 3. ft b = 3. ft H = 3. ft
	TII	PARAMETER ESTIMATES: K = 9.087E-05 ft/sec y0 = 2.925 ft
	: :	
0.001	800.	
		AQTESOLU

company: Parsons Engineering Science	риолест: 722450,14	Head Test #1)	DATA SET: AM21R.DAT 10/31/96 AQUIFER MODEL: Confined SOLUTION METHOD: Bouwer-Rice TEST DATA: H0= 5.557 ft L = 3. ft L = 3. ft L = 3. ft H = 3. ft H = 3. ft H = 5.789E-05 ft/sec y0 = 5.145 ft	
CLIENT: AFCEE	LOCATION: UST Site Beale AFB, CA	MW-2 (Rising He	Displacement (ft) 0.01 0.01 1.0 0.01 1.0 1.0 1.0	

								AOTESOLU
сонтения: Parsons Engineering Science		C		ODEL: Method: ce	. t t t t t t t t t t t t t t t t t t t	PARAMETER ESTIMATES: K = 6.113E-05 ft/sec y0 = 3.262 ft		
rsons Engine	2450.14	Test #2)	DATA SET: AMZZF.DAT 10/31/96	AQUIFER MODEL: Confined SOLUTION METHOD: Bouwer-Rice	TEST DATA: H0= 2.901 rc= 0.083 rw= 0.344 L = 3. ft b = 3. ft H = 3. ft	PARAMETER K = 6.11 y0 = 3.26		
COMPANY: Pal	PROJECT: 722450.14	Head I	E - -			: . 		
		lling		-		•	1	
	Beale AFB, CA	MW-2 (Falling	- - - -	-			320. 480 Time (sec)	
	-	F	-	-		··. · · ·	160.	
CLIENT: AFCEE	LOCATION: UST Site		10. 匠		lacement (ft)	gs:u 6 6	0.001	
CLIE	LOCA			-	lacement (ft)	qzia		

ng Science			TH SET: 31/96 JIFER MODEL: ## ined JUTION METHOD: ## ined JUTION METHOD: ## ined JUTION METHOD: ## ined JUTION METHOD: ## ined JUTION METHOD: ## ined JUTION METHOD: ## ined JUTION METHOD: ## ined JUTION METHOD: ## ined JUTION METHOD: ## ined #	AQTESOLU
сонтент: Parsons Engineering Science	рвојест: 722450.14	Head Test #2)	DATA SET: AMZZR.DAI 10/31/96 AQUIFER MODEL: Confined SOLUTION METHOD: Bouwer-Rice TEST DATA: H0= 6.98 ft r_c= 0.083 ft r_c= 0.083 ft r_d= 0.344 ft L = 3. ft L = 3. ft H = 3. ft H = 3. ft H = 3. ft DARAMETER ESTIMATES: K = 5.708E-05 ft/se y0 = 6.393 ft	
40 3	Beale AFB, CA	MW-2 (Rising He	! □	
CLIENT: AFCEE	LOCATION: UST Site		Displacement (ft) 10 10 10 10 10 10 10 10 10 1	

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APPENDIX E

MODEL INPUT AND OUTPUT FILES AND HYDRAULIC TESTING FILES

APPENDIX F

REMEDIAL ALTERNATIVE COST CALCULATIONS

Intrinsic Remediation Treatability Study UST Sites - Beale AFB, California

Present Worth Analysis

Annual Adjustment Factor = 7%

Intrinsic Remediation with Institutional		Present							
Controls and Long-Term Groundwater		Worth	Ö	ost (in curr	ent \$) at Ye	Cost (in current \$) at Year Indicated (all years not shown)	d (all years	not shown	_
and Surface Water Monitoring	years	(\$)	1	2	3	4	5	10	15
Maintain Institutional Controls	15	\$45,540	\$5,000	\$5,000	\$5,000	\$5,000	\$5,000	\$5,000	\$5,000
Long-term Monitoring									
Install New Wells	-	\$5,530	\$5,917	\$0	\$0	\$0	\$0	\$0	\$0
Groundwater/Surface Water Sampling	15	\$42,928	\$13,367	\$3,342	\$3,342	\$3,342	\$3,342	\$3,342	\$3,342
Reporting/Project Mgmt	15	\$9,883	\$1,085	\$1,085	\$1,085	\$1,085	\$1,085	\$1,085	\$1,085
Subtotal Present Worth (\$)		\$103,880						7	

Total Present Worth Cost (\$):

\$103,880

Long-Term Monitoring and Institutional Controls

Standard Rate Schedule

			Install New		Sampling		Reporting/PM
ling Category	Billing	Task 1	LTM	Task 2	(per event)	Task 3	(per event)
Cost Code/(Billing Category)	Rate	(hrs)	Wells (\$)	(hrs)	(\$)	(hrs)	(\$)
Word Processor 88/(15)	\$30	0	\$0	0	\$0	3	\$90
CADD Operator 58/(25)	\$47	2	\$94	0	\$0	0	\$0
Technician 42/(50)	\$40	4	\$160	24	\$960	5	\$200
Staff Level 16/(65)	\$57	16	\$912	8	\$456	8	\$456
Project Level 12/(70)	\$65	6	\$390	2	\$130	4	\$260
Senior Level 10/(80)	\$85	0	\$0	0	\$0	0.25	\$21
Principal 02/(85)	\$97	0	\$0	0	\$0	0	\$0
Total Labor (hrs \$)		28	\$1,556	34	\$1,546	20.25	\$1,027
ODCs							
Phone			\$14		\$15		S7
Photocopy			\$2		\$1		\$2
Mail			\$8		\$100		\$8
Computer			\$19		\$19		\$30
CAD			\$30		\$0		\$0
WP			\$0		\$0		\$11
Travel			\$99		\$99		\$0
Per Diem			\$200		\$200		\$0
Eqpt. & Supplies			\$50		\$102		\$0
Total ODCs			\$421		\$536		\$ 58
side Services							
LTM/POC Well Installation Cos	ts ^{a/}		\$3,700		\$0		\$0
Laboratory Fees b/		Soils	\$240	3 MWs, 3 SWs,	\$1,260		\$o
Other: Maintain Institutional Co	ontrols			and 2 qa/qc,			\$5,000
Total Outside Services			\$3,940		\$1,260		\$5,000

Proposal Estimate	Task 1	Task 2	Task 3
Labor	\$1,556	\$1,546	\$1,027
ODC's	\$421	\$536	\$58
Outside Services	\$3,940	\$1,260	\$5,000
Total by Task	\$5,917	\$3,342	\$6,085

Task 1: Install New LTM/POC Wells

^{a/} 2 Wells, 15ft @ \$70/ft, \$2000mob, \$100 soil handling

Task 2: Sampling per Event

^{b/} BTEX/TPH @ \$60ea, Electron acceptors at LTM wells @ \$75ea

Task 3: Reporting and PM per Sampling Event

Beale AFB Backup Calculations

Long-term Monitoring							
		Cost calculations					
Misc calculations		Description	Onit	Qty.	Unit Qty. Unit Price Subtotal	Subtotal	Total Source (If applicable)
Number of LTM wells:		Well Installation					\$ 3,700
Number of wells:	7	Mobilization	ea	-	\$ 2,000 \$ 2,000	\$ 2,000	
Depth each:	15 ft	Well Installation	In ft	30	\$ 50	50 \$ 1,500	
		Soil Disposal	drum	2	\$ 100 \$	\$ 200	

APPENDÎX F

REMEDIAL ALTERNATIVE COST CALCULATIONS

Intrinsic Remediation Treatability Study UST Sites - Beale AFB, California

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Annual Adjustment Factor = 7%

i caciii moitii Ailaiyaia			All lidal Adjustification and $r > r$	מפוווכווו ו	0 - 100				
Intrinsic Remediation with Institutional		Present	!				=		
Controls and Long-Term Groundwater		Worth	ŏ	ost (in curre	ent \$) at Ye	ar Indicate	Cost (in current \$) at Year Indicated (all years not shown)	not shown	_
and Surface Water Monitoring	years	(\$)	1	2	3	4	5	10	15
Maintain Institutional Controls	15	\$45,540	\$5,000	\$5,000	\$5,000 \$5,000	\$5,000	\$5,000	\$5,000	\$5,000
Long-term Monitoring									
Install New Wells	1	\$5,530	\$5,530 \$5,917	\$0	\$0	\$0	\$0	\$0	\$0
Groundwater/Surface Water Sampling	15	\$42,928	\$42,928 \$13,367	\$3,342	\$3,342	\$3,342	\$3,342	\$3,342	\$3,342
Reporting/Project Mgmt	15	\$9,883	\$1,085	\$1,085	\$1,085	\$1,085	\$1,085 \$1,085	\$1,085	\$1,085
Subtotal Present Worth (\$)		\$103,880							

Total Present Worth Cost (\$):

\$103,880

Long-Term Monitoring and Institutional Controls

tandard Rate Schedule

			install New		Sampling		Reporting/PM
Billing Category		Task 1	LTM	Task 2	(per event)	Task 3	(per event)
Cost Code/(Billing Category)	Rate	(hrs)	Wells (\$)	(hrs)	(\$)	(hrs)	(\$)
Word Processor 88/(15)	\$30	0	\$0	0	\$0	3	\$90
CADD Operator 58/(25)	\$47	2	\$94	0	\$0	0	\$0
Technician 42/(50)	\$40	4	\$160	24	\$960	5	\$200
Staff Level 16/(65)	\$57	16	\$912	8	\$456	8	\$456
Project Level 12/(70)	\$65	6	\$390	2	\$130	4	\$260
Senior Level 10/(80)	\$85	. 0	\$0	0	\$0	0.25	\$21
Principal 02/(85)	\$97	0	\$0	0	\$0	0	\$0
Total Labor (hrs \$)		28	\$1,556	34	\$1,546	20.25	\$1,027
ODCs							
Phone			\$14		\$15		\$7
Photocopy			\$2		\$1		\$2
Mail		1	\$8		\$100		\$8
Computer	0.71		\$19	/	\$19		\$30
CAD	. 91		\$30		\$0		\$0
WP	1 (2)		\$0		\$0		\$11
Travel			\$99		\$99		\$0
Per Diem			\$200		\$200		\$0
Eqpt. & Supplies			\$50		\$102		\$0
Total ODCs			\$421		\$536		\$58
Outside Services							
LTM/POC Well Installation Cos	sts a/		\$3,700		\$0		\$0
Laboratory Fees b/		Soils	•	3 MWs, 3 SWs,	\$1,260		\$0
Other: Maintain Institutional C	ontrols	00113		and 2 qa/qc,	Ψ1,200		\$5,000
Total Outside Services			\$3,940		\$1,260		\$5,000

Proposal Estimate	Task 1	Task 2	Task 3
Labor	\$1,556	\$1,546	\$1,027
ODC's	\$421	\$536	\$58
Outside Services	\$3,940	\$1,260	\$5,000
Total by Task	\$5,917	\$3,342	\$6,085

Task 1: Install New LTM/POC Wells

^{a/} 2 Wells, 15ft @ \$70/ft, \$2000mob, \$100 soil handling

Task 2: Sampling per Event

^{b/} BTEX/TPH @ \$60ea, Electron acceptors at LTM wells @ \$75ea

Task 3: Reporting and PM per Sampling Event

Beale AFB Backup Calculations

ong-term Monitoring								***
		Cost calculations						
Misc calculations		Description	Unit	Qty.	Unit Qty. Unit Price Subtotal	Subtotal		Total Source (If applicable)
Number of LTM wells:		Well Installation					\$ 3,700	
Number of wells:	2	Mobilization	ea	-	\$ 2,000	2,000 \$ 2,000		
Depth each:	15 ft	Well Installation	In ft	30	S	50 \$ 1,500		
		Soil Disposal	drum	2	\$ 100 \$	\$ 200		